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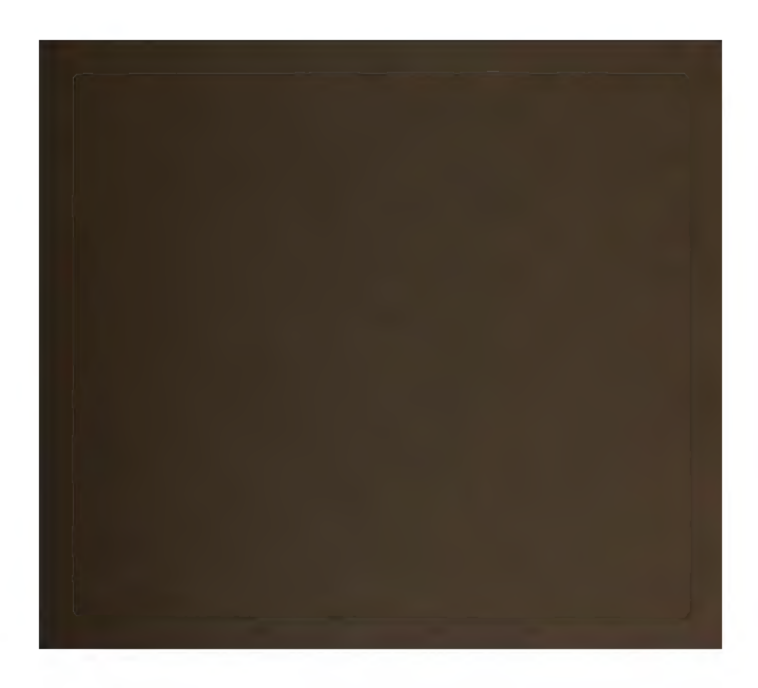
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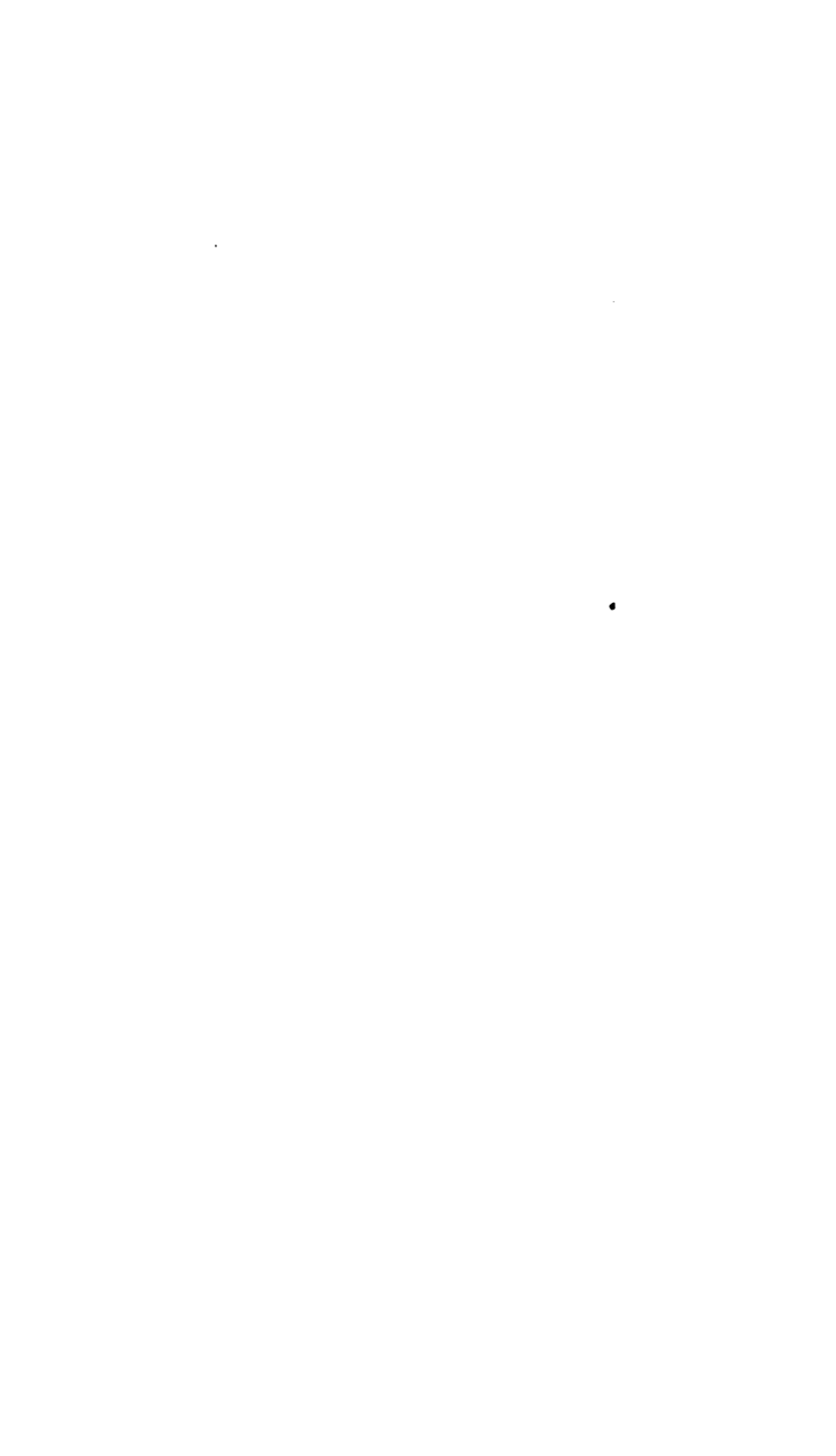
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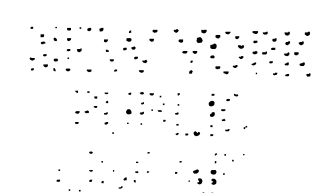
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THE FOURTH EDITION.

Vol. II.



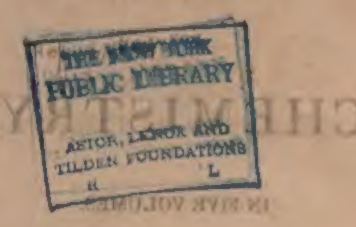
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SYSTEM

OF

CHEMISTRY;

BOOK II.

OF COMPOUND BODIES.

COMPOUND BODIES are substances composed of two or more simple bodies combined together. Now as the simple bodies described in the preceding book, excluding light and heat, are 35 in number, if they were all capable of combining together, the compounds formed by them would amount to a great many thousands: But all the simple substances are not capable of combining with each other; azote, for instance, has never been combined with metals. This diminishes their number considerably. Besides, we are still too little acquainted with the nature of caloric and light to be able to treat separately of the compounds into which they enter. Several numerous classes of compounds have been aiready described in the last Book; for the oxides, sulphurets, phosphurets, and alloys, are real Vol. 11.

Book IL

Book II.

compounds. All these circumstances render the compounds which form the subject of this Book much less numerous than might be at first supposed.

Divisions.

Compound bodies are of two kinds. Some of them are formed by the combination of two or more simple substances with each other. Thus phosphoric acid is composed of phosphorus and oxygen; and oil, of hydrogen and carbon. Others are formed by the combination of two or more compound bodies with each other. Thus phosphate of ammonia is composed of phosphoric acid and ammonia; volatile liniment, of oil and ammonia. The first of these kinds of compounds I call PRIMARY COMPOUNDS; to the second I give the name of SECONDARY COMPOUNDS. It will be very convenient to describe each of them separately.

Salifiable bases.

Besides the 35 bodies described in the preceding Book, there are a number of others lately discovered by the sagacity of Professor Davy, and which we omitted for reasons formerly specified. These, when combined with oxygen, constitute the important classess of bodies known under the names of alkalies and earths. These substances form a distinct order by themselves, and which, for that reason, may be described separately. Now these bodies (including the volatile alkali) have the property of combining with acids, and of constituting the bases of a very numerou set of bodies called salts. This induced Lavoisier to give them the appellation of salifiable bases: a name which I shall adopt, though it be somewhat exceptionable, because I can think of no other which is not equally so. This Book then shall be divided into three parts: I. SALIFIABLE BASES. II. PRIMARY COMPOUNDS. 111. SECONDARY COMPOUNDS.

DIVISION I.

OF

SALIFIABLE BASES.

In the class of salifiable bases, it is proper to include ammonia, because its properties connect it with the most important of the other substances. The salifiable bases naturally arrange themselves under the four following heads:

- 1. Volatile alkalies
- 2. Fixed alkalies.
- 3. Alkaline earths.
- 5. Earths proper.

These shall form the subject of the four following Chapters.

Book II. Division I.

CHAP. I.

OF VOLATILE ALKALIES.

Otigin.

THE word ALKALI is of Arabian origin, and was introduced into chemistry after it had been applied to a plant which still retains the name of kali. When this plant is burnt, the ashes washed in water, and the water evaporated to dryness, a white substance remains, which was called alkali: According to Albertus Magnus, who uses the word, it signifies fax amaritadinis, "the dregs of bitterness." Alkali may be obtained from other substances besides kali. Chemists gradually discovered that bodies differing from one another in several of their properties, had been confounded together under the same name. The word, in consequence, became general, and is now applied to all bodies which possess the following properties:

Properties.

- 1. A caustic taste.
- 2. Volatilized by heat.
- 3. Capable of combining with acids, and of destroying their acidity.

^{*} Thestrum Chemicum, ii. 490.

Soluble in water even when combined with carbo-

5. Capable of converting vegetable blues to green. . The alkalies at present known are three in number: 1. Ammonia: 2. Potash: 3. Soda.

The two last are called fixed alkalies, because they require a red heat to volatilize them; the first is called wolatilealkali, because it readily assumes a gaseous form, and consequently is dissipated by a very moderate degree of heat.

The fixed alkalies belong to a succeeding Chapter; the only volatile alkali at present known, namely, ammonia, shall be described in the presents

SECT. I.

OF AMMONIA.

ANMONIA can be exhibited in a state of purity only upder the form of a gas. It may be procured in the following, mattner:

1. Put into a retort a mixture of three parts of quick- Preparalime and one part of sal ammoniac in powder. Plunge the beak of the retort below the mouth of a glass jar filled with mercury, and standing inverted in a basin of mercury. Apply the heat of a lamp to the retort : a gas comes over, which displaces the mercury and fills the jar. This gas is ammonia.

Ammonia was altogether unknown to the ancients; History. the alchymists were acquainted with it, though not in

Book 11. Division 1.

and often also dissolved in water. Basil Valentine describes the method of obtaining it. It was known by the name of volatile alkali; it was also called bartsborn, because it was often obtained by distilling the horn of the hart; spirit of urine, because it may be obtained by the same process from urine; and spirit of sal ammoniac, because it may be obtained from that salt. De Black first pointed out the difference between ammonia and carbonate of ammonia, or ammonia combined with carbonic acid; and Dr Priestley discovered the method of obtaining it in a state of purity, by the process described in the beginning of this Section.

Properties.

2. Ammonia in the state of gas is transparent and colourless like air; its taste is acrid and caustic like that of the fixed alkalies, but not nearly so strong, nor does it like them corrode those animal bodies to which it is applied: its smell is remarkably pungent, though not unpleasant when sufficiently diluted. Its use as a stimulant to prevent fainting is well known.

Animals cannot breath it without death. When a lighted candle is let down into this gas, it goes out three or four times successively; but at each time the same is considerably enlarged by the addition of another same of a pale yellow colour, and at last this same descends from the top of the vessel to the bottom *.

Its specific gravity, according to the experiments of Kirwan, is 0.600, that of air being 1.00 †. While Mr Davy found it 0.5505‡. Messrs Allen and Pepys

^{*} Priestley, ii. 381.

[†] On Phlogiston, p. 28.

Chap. L

hundred cubic inches of this gas weigh, according to Kirwan, 18.26 grains, according to Davy, 17.068, according to Allen and Pepys, 18.67: Hence it is to common air nearly as 3 to 5.

When exposed to a cold of —45° it is condensed into a liquid, which again assumes the gaseous form when the temperature is raised †. When passed through a red hot tube of porcelain or glass, it is totally decomposed and converted into hydrogen and azotic gas ‡. That this experiment may succeed, the diameter of the tube must not be too great.

3. It combines very rapidly with water. When a bit of ice is brought into contact with this gas, it melts and absorbs the ammonia, while at the same time its temperature is diminished. Cold water absorbs this gas almost instantaneously, and at the same time heat is evolved, and the specific gravity of the water is diminished. Water, by my trials, is capable of absorbing 780 times its bulk of this gas; while, in the mean time, the bulk of the liquid increases from 6 to 10. specific gravity of this solution is 0.900, which just accords with the increase of bulk. It is in this state that ammonia is usually employed by chemists. term ammonia almost always means this liquid solution of ammonia in water. When heated to the temperature of about 130°, the ammonia separates under the When exposed to the temperature o form of gas.

Dany on the decomposition and composition of the fixed alkalies. Phil. Terms. 1808.

⁴ Morveau, Ann. de Chim. Ixix. 292.

prienley, ii. 305.

Book II. Division L.

-46° it crystallizes; and when suddenly cooled down, to -68°, it assumes the appearance of a thick jelly, and has scarcely any smell*.

It follows from the experiments of Mr Davy, that a saturated solution of ammonia is composed of

74.63 water 25.37 ammonia

160.00

The following TABLE, drawn up by the same ingenious chemist, exhibits the proportion of water and ammonia contained in 100 parts of liquid ammonia of different specific gravities †.

Specific gravity.	Ammonia.	Water.
0.9054	25.37	74.63
0.9166	22.07	77.93
0.9255	19.54	80.46
0.9320	17.52	82.48
0.9385	15.88	84.12
0.9445	14.53	85.47
0.9476	13.46	. 86.54
0.9513	12.40	87.60
0.9545	11.26	88.44
0.9573	10.82	89.18
0.9597	10.17	89.83
0.9619	. 8.60	90-40
0.9684	9.50	90.50
0.9030	9.09	90.01
0.9713	7.17	92.83

4. Ammoniacal gas is not altered by light; but when

Fourcroy and Vauquelin, Ann. de Chine xxix. 289.

[†] Davy, p. 68.

lectric sparks are made to pass through it, the bulk of the gas is considerably increased, and it is converted Actual of to hydrogen gas and azotic gas . Hence it follows electricity. that ammonia consists chiefly of hydrogen and azoto. By this process Berthollet converted 1.7 cubic inches of ammoniacal gas into 3.3 cubic inches +.

5. This gas has no effect upon oxygen gas while cold; Action of but when a mixture of the two gases is made to pass through a red hot porcelain tube, a detonation takes place, water is formed, and azotic gas emitted. Hence we see that ammonia is partly combustible. Its hydrocombines with the oxygen, and forms water, while the azote makes its escape in the form of a gas t. If the proportion of oxygen gas be considerable, nitric acid is also formed, in consequence of the combination of the azote with the superabundant oxygen f. The same decomposition and detonation take place if common air be used instead of oxygen gas.

orygen.

6. Sulphur is the only one of the simple combusti. Action of bles that combines with ammonia. Hydrogen pro- combustiduces no change upon it whatever; but phosphorus and charcoal act with considerable effect in high temperatures.

simple in-

It combines with sulphur in the state of vapour, and forms a sulphuret which decomposes water, and forms bydroguested sulphuret of ammonia, known formerly by the name of fuming liquor of Boyle, because it was first described by that philosopher ¶. It is commonly pre-

Pricalcy, ii. 389. + Berthollet, Jour. de Phys. 33ix. 176.

Prount, Nicholson's Journal, iii. 328. § Fourtroy, ii. 236

Shaw's Beyle, ii. 18.

Book II.
Division I.

pared by distilling a mixture of five parts of sal ammoniac, five parts of sulphur, and six of quicklime. It is a liquid of a red or rather deep orange colour, and exhales a fetid odour, in consequence of an excess of ammonia which it contains. Its nature was first pointed out by Berthollet*.

Phosphorus produces no change on ammoniacal gas while cold; but when this gas is made to pass through phosphorus in a red hot porcelain tube, it is decomposed, and phosphureted hydrogen gas, and phosphureted azotic gas are formed †.

Charcoal absorbs ammoniacal gas, but does not alter its properties while cold. But when the gas is made to pass through red hot charcoal, part of the charcoal combines with it, and forms a substance known by the name of prusic acid ‡.

Of the incombustibles,

7. Ammonia is not acted on by azote; but it combines rapidly with muriatic acid; the two gases concreting into the solid salt called muriate of ammonia.

Of metala.

8. Ammonia does not combine with the metals; but it changes some of them into oxides, and then dissolves them. The oxidizement is evidently in consequence of the decomposition of part of the water with which the ammonia is combined; for hydrogen gas is emitted during the solution. Copper and zink are oxidized by the action of ammonia; as are also tin and iron, though only superficially. Scarcely any of the other metals are altered by its action.

Liquid ammonia is capable of dissolving the oxides

[#] Ann. de Chim. xxv. 233. † Fourcroy, fl. 237, . ; Scheele, ii. 183; and Clouet, Ann. de Ellies minch a manie ?.

wilver, copper, iron, tin, nickel, zinc, bismuth, and Chap. t. cobalt . When digested upon the oxides of mercury, Dusolves lead, or manganese, it is decomposed, water is formed, by the union of the hydrogen of the ammonia with the oxygen of the oxides, and azotic gas is emitted to If a considerable heat be applied, natric acid is formed at the same time with water 1. Several other oxides are also partly deoxidized when ammonia is poured into their solutions in acids. The ammoniacal solution of the peroxide of copper is of a fine blue colour, and, according to Sage, capable of crystallizing. When heat is applied, the ammonia is partly driven off, and partly decomposed, by the combination of its hydrogen with the oxygen of the oxide.

o. Ammonia combines readily with the peroxides of gold and silver, and forms with them two compounds, formerly known by the names of fulminating gold and fulminating silver; because, when heated or rubbed, they explode with great violence. It combines also with the red oxide of mercury.

Fulminating gold, known also by the name of aurate Fulminaof ammonia, may be prepared by dissolving gold in nitro-muriatic acid, diluting the solution with thrice its weight of water, and then dropping in pure ammonia by linie and little as long as any precipitate is formed; taking care not to add too much, because in that case part of the precipitate is again dissolved. The precipi-

ting gold.

trainpolves the protozide of own when added in excess, but only in small grammaties. The peroxide of iron is involuble in it; so is the perraide of cobalt, according to Thenard.

Milner, Phil Treas.-Foureroy, v. 355.

300 to 1.

me, which is of a jeilow colour, is to be washed in pere water, dried slowly upon filtering paper, and then put into a phial: which, to prevent accidents, ought not to be corked, but its mouth covered with a linear rag or a slip of paper. This powder is fulminating gold; which is composed of five parts of yellow exide of gold and one part of ammonia. The preparation of this powder is described by Besil Valentine; and its singular properties excited the attention of all succeeding chemists. Various attempts were made to account for its falminating property, but without success, till Bergman published his dissertation on it in 1769. He demonstrated, that it is a compound of ammonia and yellow oxide of gold: that during its explosion the exide is reduced, the ammonia decomposed, and the szote, which it contained, set at liberty in the form of gas †. These facts (partly discovered by Scheele) led him to explain the explosion as follows: Ammonia is composed of azote and phlogiston. When heat is applied to fulminating gold, the phlogiston combines with the oxide, and forms gold, while the azote flies off in the form of gas. The experiments of Bergman and Scheele were repeated and confirmed by Berthollet in 1780; and the nature of exides having been previously ascertained by Lavoisier, he was enabled to give a more satisfactory explanation of the phenomenen. the explosion, the hydrogen of the ammonia combines with the oxygen of the oxide, and forms water; the gold is reduced, and the azote evolved in the form of

[•] Bergman, ü. 155.

[!] Bergman, ii. 153; and Scheele m Fire, p. 137.

ras. The great expansibility of this gas by heat ex- Chap. t. plains the violence of the explosion.

Fulminating gold explodes when struck violently, or when triturated in a mortar, or when heated to a temperature between 248° and 54°. The noise is tremendous, and when in any considerable quantity (12 grains for instance) it lacerates the metallic plate on which it is placed. When heated in close vessels, sufficiently strong to resist its action, it is reduced silently, and without any marks of violence . Its force was compated with that of gunpowder by the Royal Society, but found inferior.

Fulminating silver was discovered by Berthollet in Fulmina-1788. It may be formed by dissolving very pure silver in nitric acid, and then precipitating it by lime wazer. The precipitate is put upon filtering paper, which bsorbs the water and the nitrate of lime with which it was mixed; then pure liquid ammonia is poured upon et, and allowed to remain for 12 hours; it is then deexated off, and the black powder, on which it stood, is placed cautiously, and in very small portions, upon bits of filtering paper. This powder is fulminating silver. Even while moist it explodes with violence when struck by a hard body. When dry, the slightest touch is sufficient to cause it to fulminate. When the liquid decanted off this powder is heated in 'a glass retort, an effervescence takes place, azotic gas is emitted, and small crystals make their appearance, which are opaque, and have a metallic brilliancy. These fulminate when

[&]quot; Bergman, if. 242.

Book 11. Division 1. touched, even though covered by the liquid, and offen break in pieces the vessels in which they are kept *.

The theory of this dangerous powder is the same as that of fulminating gold. It is a compound of ammonia and oxide of silver. Friction, or the application of heat, occasions the combination of the oxygen of the oxide with the hydrogen of the ammonia; water is formed, the silver is reduced, and azotic gas emitted.

Ammoniacal fulminating mercury.

. .

Ammoniacal fulminating mercury was lately discovered by Fourcroy. It may be formed by digesting a strong solution of ammonia in water upon the red oxide of mercury. After the process has continued for eight or ten days, the oxide assumes a white colour, and is at last covered with small crystalline scales. In this state it detonates loudly upon ignited coals in the same manner as fulminating gold. It loses its fulminating property, and undergoes spontaneous decomposition, in a few days. When exposed to a low heat, the ammonia is driven off, and the red oxide assumes its former appearance †.

· Composio

As ammonia has the property of detonating with nitre, chemists had unanimously agreed that it contains phlogiston. Scheele first demonstrated, that when it is decomposed by means of the oxides of manganese, arsenic, or gold, azotic gas is set at liberty, while the oxide is reduced ‡. Hence he concluded, that it is composed of azote and phlogiston; and Bergman coincided with him in opinion. Dr Priestley discovered, that

^{*} Berthollet, Ann. de Chim 1. 54.

[†] Journals of the Royal Institution, i. 256.

^{\$} Scheele, i. 95 and 155. French Trans.—Scheele on Fire, p. 13.4

when electric explosions are made to pass through this . Chap. I. gas, its bulk is gradually augmented to thrice the space which it formerly occupied; and a quantity of hydrogen gas is produced. The same ingenious philosopher applied heat to the red oxides of mercury and lead confined in ammoniacal gas. The oxides were reduced, water was evolved, the ammoniacal gas disappeared, and instead of it, there was found a quantity of azotic gas. These experiments, and those of Scheele, led to the conclusion, that ammonia is composed of azote and hydrogen; a conclusion which was fully established by the experiments of Berthollet, published in the Memoirs of the French Academy for 1785. This acute philosopher repeated the experiments of Scheele and Priestley, and applied to them the theory of Mr Lavoisier, and added also several very decisive ones of his The most important of these is the mutual decomposition of ammonia and oxymuriatic acid. When solutions of these bodies in water are mixed together, an effervescence takes place, azote is disengaged, a quantity of water formed, and the oxymuriatic acid is converted into common muriatic acid. Now the substances mixed were ammonia and oxymuriatic acid, which is composed of oxygen and muriatic acid; the products were, muriatic acid, azote, and water, which is composed of oxygen and hydrogen. The oxygen of the water was furnished by the acid; the other products must have been furnished by the ammonia, which has disappeared. Ammonia, therefore, must be composed of azote and hydrogen. It follows from Mr Ber-

Priestley, ii. 396.

Book II. Division I. thollet's experiments, that ammonia is comp parts of azote and 29 of hydrogen *. Accc Austin's calculation it is composed of 121 p and 32 of hydrogen +. Hence 100 parts are composed of about 80 parts of azote ar drogen. The experiments of Berthollet ha farther confirmed by those made more la Davy 1, and indeed were acceded to by all unexceptionable and decisive. But the une covery by Mr Davy, that the fixed alkalies oxides, led that illustrious chemist to susp monia also might contain oxygen in so sm: tion as to escape notice. The possibility of was not to be doubted, as when ammonia. posed by means of electricity or heat, the oxy present might combine with hydrogen, a small a quantity of water as to remain invi the gases evolved. A set of experiments n pose to ascertain the point, soon convinced that oxygen is actually present in this alk pure dry ammoniacal gas was passed over 1 wire confined in a platinum tube, the iron oxydized, and a little moisture was depos experiment rendered the presence of oxyge nia somewhat probable; but it cannot be c conclusive, unless we could be certain that air was effectually excluded; which it

Berthollet determined the component parts of posing it by electricity, and exploding the new gad de Phys. xxix. 177.

[†] Phil. Tress. 178

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ar it jast.

Book IL Division L

Its base is a meal.

is placed in contact with a solution of anamouis, and the circuit completed. It gradually increases in volume, and when expanded four or five times its former bulk, becomes a soft solid. The experiment is easier, and the amalgam more permanent, if sal ammoniac slightly moistened be substituted for liquid ammonia. Mr Davy made a cavity in a piece of sal ammoniac slightly moistened, placed it on a plate of platinum attached to the positive end of the galvanic battery, put into the cavity about 50 grains of mercury, and brought in contact with the mercury a platinum wire attached to the negative end of the battery. A strong effervescence took place, much heat was evolved, and the globule in a few minutes had enlarged to five times its original bulk, and had the appearance of an amalgam of zinc. This amalgam, at the temperature of 70° or 80°, is a soft solid of the coasistence of butter; at 32° it is a firm crystallized massin which small facets appear, but having no perfectly defined form. Its specific gravity is below 3. When exposed to air it soon becomes covered with a white crust of carbonate of ammonia. When thrown into water, a quantity of hydrogen is evolved equal to half its bulk, the mercury is revived, and the water becomes a weak solution of ammonia. When confined in a given portion of air, the air increases considerably in volume, and pure mercury re-appears. Ammoniacal gas amounting to 12 or 12, the volume of the amalgam is evolved, and a quantity of exygen equal to ith or ith of the ammonia disappears. When thrown into muriatic acid gas it becomes coated with muriate of ammonia, and a little hydrogen is disengaged. In sulphuric acid it becomes costed with sulphate of ammonia and sulphur. All attempts made by Mr Davy to preserve this amaltirely of water. When put into a glass tube, or when confined under naphtha or oils, the mercury separated, ammonia was formed, and a quantity of hydrogen evolved.



It is obvious, from the preceding observations, that the amalgam thus formed, consisted of the basis of ammonta combined with mercury. This basis has so strong an affinity for oxygen that it immediately decomposes water, and is converted into ammonia while hydrogen is evolved. Hence appears to be the reason why hydrogen always appears during the destruction of the amaigam, and why the amalgam cannot be preserved. The quantity of basis of ammonia contained in 50 grains of mercury thus converted into a solid amalgam, it is evident from the preceding detail cannot exceed thath of grain. That so minute a portion of matter should make so striking a change in so great a quantity of mercury, and reduce its specific gravity so enormously, is perhaps the most extraordinary fact that has yet appeared in chemistry.

Oxygen then appears to be a constituent part of ammonia; the other ingredients of it, when deprived of that principal, acquire the property of amalgamating with mercury, and of course must be of a metallic nature. This is another unexpected and extraordinary fact. What is this metal evolved from ammonia, and to which the name of ammonium has been given? Is it in fact composed of one or of two metals? Are axote

Davy's Electrochemical Researches on the Decomposition of the Earths,

Book II. Division I. and bydrogen each metals in the gaseous state, or are they metallic oxides, or do they constitute a single metal when united together? These are questions which the present state of the subject does not enable us to resolve. The experiments seem rather more favourable to the last supposition, which is the opinion entertained by Berzelius, but they are by no means capable of deciding it.

Formation
of ammonia.

Several successful attempts have been made to form ammonia artificially. Dr Austin indeed failed in his attempts to form it by uniting together hydrogen and azotic gases by means of hear, electricity, and cold. And now that we know that this alkali is of a more compound nature than this philosopher supposed, we cannot be surprized at his failure. It could not be doubted, however, that the alkali is often formed during different chemical processes. Dr Priestley and Mt Kirwan † had actually produced it even before its composition was known. It had been found, that when tin is moistened with nitric acid, and after being allowed to digest for a minute or two, a little potash or lime is added, ammonia is immediately exhaled. The nitric acid and the water which it contains are decomposed; the oxygen of each unites with the tin, and reduces it to the state of an oxide; while at the same time the hydrogen of the water combines with the azote of the acid and with some oxygen, and forms ammonia, which is driven off by the stronger affinity of the potash or lime. Dr Austin succeeded also in forming ammonia by several other methods. He introduced into a glass tube

⁴ On Air, ii. 41.

AMMONIA.

filled with mercury a little azotic gas, and then put into the gas some iron filings moistened with water. The iron decomposes the water, and combines with its oxygen; and the hydrogen, meeting with azote at the moment of its admission, combines with it, and forms ammonia. This experiment shows, that the gaseous state of the azote does not prevent the formation of ammonia. The experiment succeeded also when common air was used instead of azote, but a longer time elapsed before the extrication of ammonia became sensible. Hence it is likely that ammonia is evolved whenever iron rusts in contact with water and air *.

* Pbil. Trans. 1788, p. 379.

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CHAP. II.

OF FIXED ALKALIES.

THE fixed alkalies are distinguished from the volatile in not being gaseous. They may be exhibited pure in a solid state. Their taste is much more acrid. Two fixed alkalies only are at present known; namely, potasb and soda. They form the subject of the following Sections.

SECT. I.

OF POTASH.

Method of procuring potach. If a sufficient quantity of wood be burnt to ashes, and these ashes be afterwards washed repeatedly with water till it comes off free from any taste, and if this liquid be filtrated and evaporated to dryness, the substance which remains behind is potash; not, however, in a state of purity, for it is contaminated with several other substances, but sufficiently pure to exhibit many of its properties. In this state it occurs in commerce under the name of potash. When heated to redness, many of its

impurities are burnt off: it becomes much whiter than Chap. 11. before, and is then known in commerce by the name of pearl-asb. Still, however, it is contaminated with many foreign bodies, and is itself combined with carbonic acid gas, which blunts all its properties. It may be obtained perfectly pure by the following process:

1. Mix it with twice its weight of quicklime, and ten times its weight of pure water: Boil the mixture for some hours in a clean iron vessel, or allow it to remain for 48 hours in a close glass vessel, shaking it occasionally. Then pass it through a filter. Boil the liquid obtained in a silver vessel very rapidly, till it is so much concentrated as to assume when cold the consistence of honey. Then pour upon it a quantity of alcohol equal in weight to one-third of the pearl-ash employed. Shake the mixture, put it on the fire, let it boil for a minute or two, then pour it into a glass vessel and cork it up. The solution gradually separates itself into two strata: the lowest consists of the impurities, partly dissolved in water and partly in a solid state; the uppermost consists of the pure potash dissolved in alcohol, and is of a reddish-brown colour. Decant this alcohol solution into a silver basin, and evaporate it rapidly till a crust forms on the surface, and the liquid below acquires such consistence as to become solid on cooling. Then pour the solution into a porcelain vessel. When cold, it concretes into a fine white substance, which is pure potash. It must be broken to pieces, and put into an air-tight phial.

For this process we are indebted to Berthollet ". The

[·] Jour d. Phys. zzvili, 401.

following, which was first proposed by Lowitz of Pe+ tersburgh, is less expensive. The potash of commerce and quicklime are to be boiled together, as above described. The filtered liquor is then to be evaporated till a thick pellicle appears on its surface, and afterwards allowed to gool; and all the crystals which have formed are to be separated, for they consist of foreign salts. The evaporation is then to be continued in an iron pot; and, during the process, the pellicle which forms on the surface is to be carefully taken off with an iron skimmer. When no more pellicle ap ears, and when the matter ceases to boil, it is to be taken off the fire, and must be constantly agitated with an iron spatula while cooling. It is then to be dissolved in double its own weight of cold water. This solution is to be filtered and evaporated in a glass retort * till it begins to deposite regular crystals. If the mass consolidates ever so little by cooling, a small quantity of water is to be added, and it must be heated again. When a sufficient number of crystals have been formed, the liquor which swins over them, and which has assumed a very brown colour, must be decanted off, and kept in a well-closed bottle till the brown matter has subsided, and then it may be evaporated as before, and more crystals obtained +.

The theory of these processes is obvious: The lime separates the carbonic acid, for which it has a stronger

Dr Kennedy observes, very justly, that a glass retort ought not to be employed, because potash in this state dissolves glass. Edin. Trans. v. 97.

[†] Nicholson's Journal, i. 164.

affinity; and the sloohol or the evaporation separate all the other foreign ingredients.

"Chap. II.

A still simpler method is employed b Klaproth. He boils equal parts of salt of tartar (carbonate of potash | repared from tartar), and carrara marble or oyster shells, burnt to lime, with a sufficient quantity of water, in a polished iron kettle. The ley is then strained through clean linen, and though still turbid, is reduced by boiling, till it contain about one half its weight of potash. It is then passed a second time through a linen cloth, and set bye in a glass bottle. After some days, when the ley has become clear of itself, it is decanted off from the sediment into another bottle ".

As potash is never obtained at first in a state of puri- Black's disty, but always combined with carbonic acid, it was long the cause of before chemists understood to what the changes produced upon it by lime were owing. According to some, it was deprived of a quantity of mucilage, in which it had formerly been enveloped; while, according to others, it was rendered more active by being more comminuted. At last, in 1756, Dr Black proved, by the most ingenious and satisfactory analysis, that the potasb which the world had considered as a simple substance, was really a compound, consisting of potash and carbonic acid; that lime deprived it of this acid; and that it became more active by becoming more simple.

While Dr Black was thus occupied in Scotland, Mr Meyer was employed in Germany in the same researches; from which, however, he drew very different conclusions. His Essays on lime appeared in 1764.

covery of caunicity.

[·] Klaproth' Butrage, i, Preface, p. 10.

Pouring into lime-water a solution of potash (carbonate of potasb), he obtained a precipitate, which he found not to differ from lime-stone. The alkali had therefore deprived the lime of its causticity and its active properties; and these very properties it had itself acquired. From which he concluded, that the causticity of lime was owing to a particular acid with which it had combined during its calcination. The alkali deprived the liene of this acid, and therefore had a stronger affinity for it. To this acid he gave the name of acidum pingue or causticum. It was, according to him, a subtile clastic mixt, analogous to sulphur, approaching very nearly to the nature of fire, and actually composed of an acid principle and fire. It was expansible, compressible, volatile, astringent, capable of penetrating all vessels, and was the cause of causticity in lime, alkalies, and metals. This theory was exceedingly ingenious, and it was supported by a vast number of new and important facts. But notwithstanding the reputation and acknowledged genius and merit of its author, it never gained many followers; because the true theory of causticity, which had been already published by Dr Black, soon became known on the continent; and notwithstanding some opposition at first, soon carried conviction into every unprejudiced mind. Mr Jacquin, botanical professor at Vienna, published a latin dissertation in defence of Dr Black's doctrine in 1769. This work was opposed in 1770 by Crans, physician to the king of Prussia, who endeavoured to defend the hypothesis of Meyer, who was now dead, in a very elaborate treatise. The subject was resumed by Mr Lavoisier in 1774 in his physical and chemical essays. He repeated the experiments of Dr Black and his disciples,

and confirmed them in every particular. Since that rime the hypothesis of Meyer seems to have been abandoned by every one.

That potash was known to the ancient Gauls and Germans cannot be doubted, as they were the inventors of soap, which, Pliny informs us, they composed of ashes and tallow. These ashes (for he mentions the ashes of the beech-tree particularly) were nothing else but potash; not, however, in a state of purity . The me, too, mentioned by Aristophanes and Plato, appears to have been a ley made of the same kind of ashes. The alchymists were well acquainted with it; and it has been in every period very much employed in chemical researches. It may be said, however, with justice, that till Berthollet published his process in the year 1786, chemists had never examined potash in a state of complete purity +.

2. Potash is a brittle substance of a white colour, and Properties a smell resembling that which is perceived during the lacking of quicklime. Its taste is remarkably acrid; and it is so exceedingly corresive, that when applied to any part of the body, it destroys it almost instantaneous ly. On account of this property, it has been called

of potasb.

Plieti lib. zvii. c. 51

t Potash was long distinguished by the name of vegetable alkali, because it is obtained from vegetables, and because it was long thought to be pecular to the vegetable kingdom; but this is now known to be a mintake. It was called also solt of tartar, because it may be obtained by burning the salt called tartar. Mr Kirwan has given it the name of turters; Dr Pearson has called it organials, Klaproth iais, and Dr Black diarece By most Entish chemists tis called posoib, but this se m, in common language, ugnifies the carbonate of potash, or the potash of otherer.

caustic, and is often used by surgeons, under the name of the potential cautery, to open abscesses, and to destroy useless or hurtful excrescences. Its specific gravity is 1.7085 ‡.

When heated it melts; at a red heat it swells, and evaporates slowly in a white acrid smoke. A strong heat gives it a greenish tinge, but produces no other alteration in it. Potash is not altered by exposure to light.

When exposed to the air, it soon attracts moisture, and is converted into a liquid; at the same time it combines with carbonic acid, for which it has a strong affinity.

Its combination with water.

3. It has a very strong affinity for water. At the common temperature of the air, one part of water dissolves two parts of potash. The solution is transparent, very dense, and almost of the consistence of oil. It is in this state that potash is usually employed by chemists. When four parts of potash in powder and one of snow are mixed together, the mixture becomes liquid, and at the same time affords a quantity of caloric. This mixture was employed by Lowitz to produce artificial cold. When the aqueous solution of potash is evaporated to a proper consistency, the potash crystallizes. The shape of its crystals is very different, according to the way in which they have been produced. When allowed to form spontaneously, they are octahedrons in groupes, and contain 0.43 of water *: When formed by evaporation

Hassenfratz, Ann. de Chim. xxviii. ex.

[&]quot;According to Proust, the hydrate of potash contains only 0.30 of war ter. Jour. de Phys. lix. 266.

on the fire, they assume the figure of very thin transparent blades of extraordinary magnitude, which, by an assemblage of lines crossing each other in prodigious numbers, present an aggregate of cells or cavities, commonly so very close, that the vessel may be inverted without losing one drop of the liquid which it conteins t.

4. Potash shows no disposition to unite with oxygen, Action on acither is it altered by the action of any of the compounds into which oxygen enters, though it has a strong tendency to unite with several of these compounds.

5. It unites with none of the simple combustibles ex- Op simple cept sulphur. Carbon and hydrogen do not act upon it bles. at all; neither does it produce any alteration in them, but it acts upon phosphorus with considerable energy.

When three parts of sulphur and one of potash are Sulphuret triturated together in a glass mortar, the sulphur acquires a green colour, the mixture becomes hot, and exhales an alliaceous odour. It gradually attracts moisture from the air, and is totally soluble in water ". two parts of potash and one of sulphur are heated in a crucible, they melt and combine, and form a sulphuret of potarb. The potash of commerce may be also employed; for the carbonic acid separates in the form of a gas during the combination of the potash and sulphur. When the fusion is complete, the sulphuret is to be poured upon a marble slab; and as soon as it congeals, it must be broken in pieces, and set by in a wellcorked phial.

Sulphuret of potash, thus prepared, is of a brown co-

Intersise the liver of animals. Hence it was need depar sulphuris, "liver of sulphur;" has a subsect to the air, it soon becomes green, wake. It is hard, brittle, and has a glassy wake. Its taste is acrid, caustic, and bitter, and it wown stain upon the skin. It has no other that of sublimed sulphur. When exposed a moient hear, the sulphur sublimes, and the potash restable blues to green, and soon destroys them. When rested with charcoal, it dissolves and combines with

When sulphuret of potash is exposed to the air, or

when it is moistened with water, its properties very

the odour of sulphureted hydrogen gas. This change is owing to the formation of a quantity of sulphureted hydrogen, in consequence of the decomposition of the water. This new-formed substance combines with the sulphuret, and converts it into bydrogureted sulphuret of potash, which is soluble in water, and has a brownish green colour. It may be formed also by boiling in water two parts of potash and one part of sulphur. Sulphuret of potash produces no change upon air, but hy-

drogureted sulphuret gradually absorbs oxygen. When

inclosed in a vessel with a quantity of air, it soon ab-

sorbs all the oxygen of that portion, and leaves nothing

but azotic gas. This fact, which was first observed by

Scheele, induced him to use hydrogureted sulphuret to

measure the quantity of oxygen contained in any given

Hydrogureted sul-

[#] Fourcroy, ii. 203.

portion of atmospheric air. Hydrogueted sulphurou is . Chap. H. capable of oxidizing and dissolving almost all the metals. We are indebted to Mr Berthollet for the first accurate account of the difference between these two substances +.

Potash causes be combined with phosphorus by any method at present known. But when potash, dissolved phosphorus. in water, is heated over phosphorus in a retort, the water is gradually decomposed, part of the phosphorus is converted into phosphoric acid, and a great quantity of phosphureted hydrogen gas is emitted, which takes fire, as usual, as soon as it comes in contact with the air of the atmosphere. It was by this process that Gengembee first obtained phosphureted hydrogen gas.

Action on

6. It does not appear that potash is capable of uniting Action on with azote, or even of acting on it at all: but with combustmuriatic acid it unites very readily, and forms the compound known by the name of muriate of potash.

7. Potash does not combine with any of the metals; On metals, but some of the metals which have a strong affinity for oxygen, when put into a solution of potash in water, especially if heat be applied, are gradually oxidized. This is the case with molybdenum, zinc, and iron. Tin alis oxidized in a very small proportion; and this scems also to be the case with manganese.

It is capable of dissolving a considerable number of And theat the metallic oxides; and in some cases it deprives them of a dose of their oxygen. Thus when poured upon the red oxide of iron, it soon converts it into the black. The cause of this change is unknown. It has been as-

[&]quot; Aus. de Chiu. Exv. 233. See also Proust, Jear. de Phys. lix 265

Book If. Devision I. certained, that the oxides of the following metals are soluble in potash:

Lead*,
Tin,
Nickel,
Arsenic,
Cobalt,
Manganese,

Zinc,
Anumony,
Tellurium,
Tungsten,
Molybdenum.

But the nature of these solutions has not hitherto been examined with any degree of attention; though the subject is remarkably curious, and promises to throw light both upon the nature of alkalies and metals.

Whether a

Various opinions have been entertained by chemists respecting the composition of potash. At one time it was fashionable to consider azote as one of its constituents; because that principle had been detected in ammonia, and it was thought not unlikely by Lavoisier. that as oxygen, one of the constituents of air, was the ecidifying principle, azote, the other constituent, would be found to be the principle of alkalisation. Morveau and Desormes announced, some time ago, that they consider potash as a compound of hydrogen and lime. Their chief proofs were the appearance of lime, when the salt, composed of hyperoxygenized muriatic acid and potash, is strongly heated with phosphoric acid in a crucible of platmum; and a manifest combustion, together with the deposition of lime, when charcoal and potash are in like manner exposed to a strong heat, in a

Bergman, ili. 456. Proust, Jour. de Phys. Ivi. 207.

platinum crucible . But these and the other experiment tal proofs being examined by Darracq, that accurate chemist ascertained that the results obtained by Desormes and Morveau were owing, in most cases, to the impurity of the potash with which they had made their experiments; while in others they had drawn wrong inferences from mistaken resemblances †. Their hypothesis of course cannot be maintained.

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This question, inferior to none in the annals of che- Composimistry, has been at last decided by the happy sagacity ash. of Mr Davy, who has recently enriched chemistry with a long train of the most important and brilliant discoveries. Potash, it follows from his experiments, is a compound of exygen and a new metal, to which he has given the name of potassium.

When potash is perfectly dry it is a non-conductor of How doelectricity, but it becomes a conductor when slightly moistened on the surface, a degree of moisture which it acquires by being exposed for a few seconds to the aumosphere. When pieces of potash in this state are placed upon a disc of platinum attached to the negagive end of a powerful galvanic battery, and a platinum wire from the positive extremity is made to touch its upper surface, the potash is gradually decomposed, oxygen gas separating at the extremity of the positive wire, while globules of a white metal like mercury appear at the side in contact with the platinum disc. A number of accurate experiments demonstrated to Mr. Davy, that these globules were the basis of potash, and that they were converted into potash by absorbing

composed.

Book II.
Division I.
Properties
of potassium.

oxygen. This metallic substance possesses the follow-ing properties:

Its colour is white like that of mercury. At the temperature of 100° it is as fluid as mercury; at 60° it still continues imperfectly fluid; at 50° it is a soft and malleable solid, while at 32° it is hard, brittle, and crystallized in facets. When heated nearly to redness it is volatilized and converted into vapour. It conducts electricity and heat as well as other metallic bodies.

Its specific gravity.

It differs remarkably from all the metals previously known in its specific gravity; being lighter than any other liquid substance hitherto examined, swimming even in naphtha of the specific gravity 0.770. As it alters very rapidly when exposed to the air, it is difficult to ascertain its specific gravity with accuracy. Mr Davy endeavoured to estimate the relative weight of a globule of potassium and of mercury, of exactly the same bulk, measured by means of a micrometer: he found the weights to each other as 10 to 223, which gives 0.6 nearly for the specific gravity of potassium.

Action of oxygen.

Its affinity for oxygen and its tendency to absorb that principle are much greater than that of any other substance previously known. Hence, if exposed to the air, it absorbs oxygen, and is covered with a crust of potash in a few minutes; this crust absorbs water which is rapidly decomposed, and in a short time the whole becomes a saturated solution of potash. When thrown upon water it decomposes that liquid with rapidity, heat is evolved, hydrogen gas holding potassium in solution is emitted, which takes fire spontaneously, and the whole potassium burns with an explosion, and is converted into potash. It burns equally when placed upon ice. It equally decomposes water, and is converted in-

to potash when kept in alcohol, ether, and oils, or when Chap IL. thrown into the mineral acids. Newly distilled naphthat is the substance in which it may be best preserved. In this liquid it remains unaltered for many days, and may be readily examined in the open air when covered with a crust of it.

When heated in hydrogen gas, a portion of it is dissolved, and the gas acquires the property of burning spontaneously when it is mixed with atmospheric air. But if kept for a short time it again deposites the potassium, and loses the property of burning spontaneously.

When heated in a small quantity of oxygen gas it ab- Protoxide sorbs a portion of it, loses its metallic appearance, as- of potas sumes a reddish brown colour, and becomes grey when cold. In this state it may be considered as a protoxide of potassium. This protoxide may be readily formed by fusing together potash and potassium in a glass tube filled with the vapour of naphtha. When exposed to the air it readily absorbs more oxygen, and is converted into potash.

When potassium is exposed to the action of oxymuristic acid gas it takes fire, and is converted into muriate of potash.

Potassium, when brought in contact with phosphorus, Phosphuret. and pressed upon under naphtha, combines with it and forms a phosphuret of patassium, which has the colour of lead, and remains solid at the boiling point of naphtha. When exposed to the air it slowly absorbs oxygen and is converted into phosphate of potash. When the metal and phosphorus are brought into contact in the open air, they become fluid together, burn, and are converted at once into phosphate of potash.

:.:

Book II. Division I. Sulphuret.

Potassium combines rapidly with sulphur in tubes filled with the vapour of naphtha, while heat and light are evolved. The sulphuret formed has the grey colour of artificial sulphuret of iron. A little sulphureted hydrogen is given out during the formation of this compound. In the open air the combination takes place with combustion, and sulphuret of potash is formed. Sulphuret of potassium, when exposed to the air, is gradually converted into sulphate of potash.

Alloys

Potassium readily combines and forms an alloy with all the metals hitherto tried. When one part of potassium is added to about 10 parts of mercury in bulk, they instantly unite and form a substance very like mercury in colour. When a globule is made to touch a globule of mercury twice as bulky, they combine with considerable heat. The globule is at first fluid, but on cooling becomes solid and resembles silver. If the potassium be increased to about to the mercury in weight, the amalgam is harder and becomes brittle. . One part of potassium renders 70 parts of mercury solid, and forms an amalgam very soft and malleable. When these amalgams are exposed to the air, they rapidly absorb oxygen, potash is formed, which deliquesces, and in a few minutes the mercury is found pure and unaltered. In water the amalgam is decomposed with a hissing noise, hydrogen gas is evolved, and the mercury remains free. The fluid a lgam of potassium acts upon most metals. In this state of union mercury acts on iron and platinum.

When potassium is heated with gold, silver, or copper in a close glass vessel, it combines with them. The alloy is decomposed when thrown into water, potash is formed, and the other metals separated. No attempts

have been made to combine it with other metals, except with fusible metal, with which it unites and form's an alloy that requires more heat to melt it than the fusible metal itself.

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When potassium is mixed and heated with metallic Action on oxides, it rapidly reduces them to the metallic state. Mr ides. Davy tried the experiment on the oxides of iron, tin, and lead. When there is an excess of potassium it forms an alloy with the revived metal. Its action upon metallic oxides is so strong, that it readily decomposes flint and green glass in a gentle heat.

Mr Davy demonstrated by the most decisive expe- Constituriments, that when potassium combines with oxygen to saturation it is converted into potash. He performed this experiment in glass tubes, and ascertained the bulk of oxygen gas absorbed during the combustion of a given From these experiments it folweight of potassium. lows, that potash is composed of about 6 parts potassium and 1 part of oxygen, or nearly of

ents of pot-

potassium 86

oxygen

100

Galvanism is not the only mode by which potash may Potash debe decomposed, and its base obtained nearly in a state by iron. of purity. Gay Lussac and Thenard have succeeded in decomposing it by means of iron filings; and their experiment has been successfully repeated by Mr Davy. Into the middle of an iron gun barrel is to be put a quantity of clean and dry iron filings or turnings. An iron tube, containing potash as dry as possible, should be ground to one end of the gun barrel, and having a

compose

small hole through which the potash may run slowly when melted. To the other extremity a tube of safety, containing mercury or naphtha, ought also to be luted, and great care should be taken that all the lutings be air tight. The gun barrel being laid across a furnace, the iron turnings within it are brought to a white heat, while the potash is kept cool by means of ice; then the potash is brought into fusion, and made to flow slowly through the iron turnings. Hydrogen gas is emitted in considerable quantity during the whole process. The part of the gun barrel next the tube of safety should be kept constantly cool. When the process is at an end, a portion of potassium nearly pure is found near the tube safety; but the greatest part of it is alloyed with the iron turnings.

Such are the properties of potassium hitherto investigated. For all the facts above stated we are indebted to
Mr Davy +; but his experiments have been repeated
and confirmed by other respectable philosophers. Mr
Davy has lately ascertained that the protoxide of potassium has the property of combining with ammonia and
azote. The last compound inflames spontaneously
when exposed to the air, potash is formed, and azotic
gas disengaged. It acts violently on water, and produces potash and ammonia ‡.

Potash or the peroxide of potassium is of the highest

[•] Phil. Mag. 222ii. 89, and 276. See also a note in Mr Davy's paper Cu the Decomposition of the Earths. Phil. Trunt. 1808.

[†] See Davy's lecture on the Decempesition and Composition of the Fined

^{\$} Phil. Mag. 22211. 368

importance, not only in chemistry, where it is employed for a great variety of purposes, but also in many arts and manufactures; as washing, bleaching, dyeing, glass-making, and others, as will afterwards appear. It is employed also in surgery and medicine.

SECT. II.

OF SODA.

Soda, called also fossil or mineral alkali*, because it Names, was thought peculiar to the mineral kingdom, was known to the ancients (though not in a state of purity) under the names of vileov and nitrum +.

It is found in large quantities combined with carbo- Preparanic acid in different parts of the earth, especially in Egypt; and common salt is a compound of soda and muriatic acid. But the soda of commerce is obtained from the ashes of different species of the salsola, a genus of plants which grow upon the sea-shore, especially from the salsola soda, from which the alkali has obtained its name. The soda of commerce is also called barilla, because the plant from which it is obtained bears that name in Spain. Almost all the algæ, especially the

Dr Pearson has proposed to distinguish it by the name of fossalkali; Klaproth calls it natron.

[†] The 2.7pov of the Athenians was evidently the same substance; and so was the and of the Hebrews.

Book 11. Division 1. fuci, contain also a considerable quantity of soda. The ashes of these plants are known in this country by the name of kelp; in France they are called varec.

The soda, or barilla, of commerce, is far from being pure; besides carbonic acid it contains common salt, and several other foreign ingredients; but it may be obtained perfectly pure by the processes described in the last Section for purifying potash.

Soda and potash resemble each other so nearly, that they were confounded together till Du Hamel published his dissertation on common salt in the Memoirs of the French Academy for 1736. He first proved that the base of common salt is soda, and that soda is different from potash. His conclusions were objected to by Pott, but finally confirmed by Margraff in 1758.

Properties.

Soda is of a greyish-white colour, and agrees exactly with potash in its taste, smell, and action upon animal bodies; but its specific gravity is only 1.336 †.

Heat produces on it exactly the same effects as upon potash. When exposed to the air, it absorbs moisture and carbonic acid, and is soon reduced to the consistence of paste: but it does not liquefy like potash; in a few days it becomes dry again, and crumbles into powder.

It has a strong affinity for water, dissolves in it like potash, and may also be obtained in crystals by evaporating its aqueous solution. It is not altered by light; nor does it combine with oxygen, hydrogen, azote, carbon, charcoal, nor metals. Its action upon phosphorus and sulphur is the same with that of potash. The sul-

^{*} Opusc. ii. 331.

[†] Hassenfratz, Ann. de Chim. xxviii. II.

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phuret and hydrogureted sulphuret of sods possess the properties of the sulphuret and hydrogureted sulphuret of potash, and are formed in the same manner. In its oction on metals, metallic oxides, and in its affinities, it also agrees with potash. In short, the two fixed alkalies, in a state of purity, resemble each other very nearly in almost every particular.

Similar opinions respecting its composition were en- Compositertained by chemists, as those which they had respecting the composition of potash. Fourtroy supposed it a compound of magnesia and azote. Desormes and Morreau*, on the other hand, affirmed that it is composed of magnesia and hydrogen; but the experiments upon which this opinion was founded have been proved inoccurate by Darracq t. Mr Davy has lately succeeded in decomposing it by the same processes which enabled him to ascertain the composition of potash. Like that alkali it is a metallic peroxide. To its metallic basis Mr Davy has given the name of sodium.

Soda is decomposed by the galvanic battery in the ame way as potash; but requires a more powerful battery, or much smaller pieces of soda must be exposed to its action.

Sodium is a white metal like silver, and at the com- Properties mon temperature of the atmosphere is solid; but exceedingly malleable, and so soft that pieces of it may be welded together by strong pressure. It still retains its malleability and softness at the temperature of 32°. When heated to 120° it begins to melt, and is completely fluid at 180°; though exposed to a red heat

strong enough to melt plate glass, none of it is volatilized.

It conducts electricity and heat in the same manner as potassium. It is heavier than that metal, though not so heavy as water; swimming in oil of sassafras, of the specific gravity 1.096, and sinking in naphtha of the specific gravity 0.861. By mixing these two liquids together till they acquired just the specific gravity of sodium, Mr Davy ascertained that the specific gravity of that metal is 0.9348.

Its affinity for oxygen is similar to that of potassium, and hence it acts nearly in a similar manner when exposed to substances containing that principle. When exposed to the air it absorbs oxygen, and is soon covered with a crust of soda, which deliquesces much more slowly than potash; hence the sodium is not so soon converted into an alkali as potassium. No combustion takes place unless the sodium is heated nearly to redness; but the rapidity of the absorption of oxygen increases with the temperature. The flame which it produces in oxygen gas is white, and as it sends out bright sparks the effect is beautiful.

Hydrogen gas though assisted by heat seems to have no action on it whatever. When thrown into water so-dium occasions a violent effervescence, with a loud hissing noise; hydrogen gas is evolved and soda formed; but no combustion takes place as happens to potassium, probably because sodium is insoluble in hydrogen gas. A few scintillations indeed appear when sodium is thrown into hot water. Sodium usually burns also when brought in contact with a small particle of water, or when placed on moistened paper.

Protozide.

When fused with dry soda in certain quantities, there

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is a division of oxygen between the sods and the base; and a protoxide of sodium is formed of a deep brown colour while fluid, but which becomes a dark grey solid on cooling. This protoxide when exposed to the air absorbs oxygen, and is converted into soda.

When sodium is exposed to the action of oxymutiatic acid gas, it burns vividly with bright scintillations of a red colour.

It combines with phosphorus with the same pheno- Phosphumena as potassium, and forms a similar phosphuret, which is converted into phosphate of soda by exposure to the air.

It combines with sulphur in close vessels filled with Sulphures. the vapour of naphtha with great vividness, with light and heat, and often with explosion from the vaporization of a portion of the sulphur, and the disengagement of sulphureted hydrogen. The sulphuret of sodium is of a deep grey colour.

It decomposes the water in alcohol and ether precisely as potassium does. In oils it gradually absorbs oxygen and forms soaps. It is converted into soda when thrown into the mineral acids, in nitric acid with infammation, and in sulphuric and muriatic acids with the evolution of much heat.

It combines with metals, and forms alloys similar to Alloys the alloys of potassium. One part of it renders 40 parts of mercury solid, and of the colour of silver; and the combination is attended with a considerable degree of heat. It combines with tin without changing its colour, and acts on gold and lead by the assistance of heat. When these alloys are exposed to the air the sodium soon absorbs oxygen, and is converted into sode;

Composition of soda. the amalgam of sodium combines with the other metals and with sulphur, forming triple compounds.

From a number of experiments on the combination of sodium with oxygen, made in the same manner as those on the combination of potassium with the same principle, Mr Davy has shown that soda is composed of about 7 parts sodium and 2 of oxygen, or nearly of

Sodium 78

Oxygen 22

100

Thus it appears that both the fixed alkalies are metallic peroxides, and that the bases of both are capable of combining with two doses of oxygen. But the protoxide is not permanent in consequence of its great affinity for oxygen.

The importance of soda in manufactures is not inferior to that of potash. For several purposes, indeed, as for the manufacture of soap and glass, it answers even better than potash.

CHAP. III.

OF THE ALKALINE EARTHS.

THE word EARTH, in common language, has two meanings; it sometimes signifies the globe, which we inhabit, and sometimes the mould on which vegetables grow. Chemists have examined this mould, and have found that it consists of a variety of substances mixed together without order or regularity. The greatest part of it, however, as well as of the stones which form ²pparently so large a proportion of the globe, consists of a small number of bodies, which have a variety of common properties. These bodies chemists have agreed to class together, and to denominate earths.

Every body which possesses the following properties is an earth.

1. Insoluble in water, or nearly so; or at least beco- properties. ming insoluble when combined with carbonic acid.

- 2. Little or no taste or smell; at least when combined with carbonic acid.
- 3. Fixed, incombustible, and incapable while pure of being altered by the fire.
 - 4. A specific gravity not exceeding 4.9.

- 5. When pure, capable of assuming the form of a white powder.
 - 6. Not altered when heated with combustibles.

The earths have been divided into two classes, namely, alkaline earths and earths proper. The first have the property of giving a green colour to vegetable blues, and of neutralizing acids; the second do not alter vegetable blues, and are incapable of neutralizing acids.

The alkaline earths are four in number; namely,

Lime,
Magnesia,
Barytes,
Strontian.

The properties of these bodies occupy our attention in the four following Sections.

SECT. I.

OF LIME.

LIME has been known from the earliest ages. The ancients employed it in medicine; it was the chief ingredient in their mortar; and they used it as a manure to-fertilize their fields.

Lime abounds in most parts of the world, or perhaps. I should rather say, that there is no part of the world where it does not exist. It is found purest in lime-stones, and marbles, and chalk. None of these substances, however, is, strictly speaking, lime; but they are all capable of becoming lime by a well-known pro-

cess, by keeping them for some time in a white heat : this process is called the burning of lime. The product, which in common language is denominated quicklime, is the substance known in chemistry by the name of

1. Lime may be obtained perfectly pure by burning Preparathose crystallized limestones, called calcarcous spars, which are perfectly white and transparent, and also by burning some pure white marbles. It may be procured also in a state of purity by dissolving oyster-shells in muriatic acid, filtering the solution, mixing it with ammonia as long as a white powder continues to fall, and filtering again. The liquid is now to be mixed with a solution of carbonate of soda: the powder which falls being washed and dried, and heated violently in a platinum crucible, is pure lime.

2. Pure lime is of a white colour, moderately hard, properties but easily reduced to a powder.

It has a hot burning taste, and in some measure corrodes and destroys the texture of those animal bodies to which it is applied. Its specific gravity is 2.3. It tinges vegetable blues green, and at last converts them to rellow.

It is incapable of being fused by the most violent beat that can be produced in furnaces, or even by the most powerful burning-glasses.

3. If water be poured on newly burnt lime, it swells Stacking of and falls to pieces, and is soon reduced to a very fine powder. In the mean time, so much heat is produced. that part of the water flies off in vapour. If the quan-

tity of lime slacked (as this process is termed) be great, the heat produced is sufficient to set fire to combusti-In this manner, vessels loaded with lime have sometimes been burnt. When great quantities of lime are slacked in a dark place, not only heat but light also is emitted, as Mr Pelletier has observed *. When slacked lime is weighed, it is found to be heavier than it was before. This additional weight is owing to the combination of part of the water with the lime; which water may be separated again by the application of a red heat; and by this process the lime becomes just what it was before being slacked +. Hence the reason of the heat evolved during the slacking of lime. of the water combines with the lime, and thus becomes solid; of course it parts with its caloric of fluidity, and probably also with a considerable quantity of caloric which exists in water even when in the state of ice: for when two parts of lime and one part of ice (each at 32°) are mixed, they combine rapidly, and their temperature is elevated to 212°. The elevation of temperature during the slacking of barytes and strontian is owing to the same cause. From the experiments of Mr Dalton it follows, that slacked lime, well dried in a moderate heat, is composed of 3 parts lime and 1 part water 1. This result does not differ much from the previous experiments of Lavoisier, who found that 1000 parts of lime, when slacked, were con-

Hydrate of lime.

[·] Jour. de Phys. i. 22.

[†] Dr Black.

¹ Dalton's New System of Chemical Philosophy, i. 87.

verted into 1287 parts ". Slacked lime then may be Chap. 111. considered as a hydrate of lime.

The smell perceived during the slacking of lime is owing to a part of that earth being elevated along with the vapour of the water; as evidently appears from this circumstance, that vegetable blues exposed to this vapour are converted into green.

Limestone and chalk, though they are capable of being converted into lime by burning, possess hardly any of the properties of that active substance. They are tasteless, scarcely soluble in water, and do not perceptibly act on animal bodies. Now, to what are the new properties of lime owing? What alteration does it undergo in the fire?

It had been long known, that limestone loses a good deal of weight by being burned or calcined. It was nafural to suppose, therefore, that something is separated from it during calcination. Accordingly, Van Helmont. Ludovicus, and Macquer, made experiments in succeseason, in order to discover what that something is; and whey concluded from them that it is pure water, which the lime recovers again when exposed to the atmosphere. As the new properties of time could hardly be ascribed to this loss, but to some other cause, Stahl's opinion. like all the other chemical theories of that wonderful man, was generally acceded to. He supposed that the new properties which lime acquired by calcination are owing entirely to the more nanute division of its particles by the action of the fire. Boyle indeed had endeavoured to prove, that these properties are owing to

Lavonice's Essays translated by Henry, p. 230.

braced by Newton and illustrated by Hales, and which Meyer new modelled, and explained with so much ingennity and acuteness as to draw the attention of the most distinguished chemists. But while Meyer was thus employed in Germany, Dr Black of Edinburgh published in 1756 those celebrated experiments which form so brilliant an era in the history of chemistry.

Discovered by Drilling

. He first ascertained, that the quantity of water separated from limestone during its calcination is not nearly equal to the weight which it lost. He concluded in consequence, that it must have lost something else than mere water. What this could be, he was at first at a loss to conceive; but recollecting that Dr Hales had proved that limestone, during its solution in acids, emits a great quantity of air, he conjectured that this might probably be what is lost during calcination. Hecalcined it accordingly, and applied a pneumatic apparatus to receive the product. He found his conjecture verified; and that the air and the water which separated from the lime were together precisely equal to the loss of weight which it had sustained. Lime therefore owes its new properties to the loss of air; and limestone differs from lime merely in being combined with a certain quantity of air: for he found that, by restoring again the same quantity of air to lime, it was converted into limestone. This air, because it existed in lime in a fixed state, he called fixed air. It was afterwards examined by Dr Priestley and other philosophers; found to possess peculiar properties, and to be that species of gas now known by the name of carbonic acid gas. Lime then is the simple substance, and limestone is composed of carbonic acid and lime. Heat separates

the carbonic acid, and leaves the lime in a state of Chap. HIL purity.

5. When lime is exposed to the open air, it gradually attracts moisture, and falls to powder; after which it soon becomes saturated with carbonic acid, and is again converted into carbonate of lime or unburnt lime-

Water, at the common temperature of the atmosphere, Action of dissolves less than 0.002 parts of its weight of lime . This solution is called lime-water. It is limped, has an acrid taste, and changes vegetable blue colours to green. One ounce troy of lime-water contains about one grain of lime. It is usually formed by throwing a quantity of lime in powder into pure water, allowing it to remain for some time in a close vessel, and then decanting the transparent solution from the undissolved lime. When lime-water is exposed to the air, a stony crust soon forms on its surface composed of carbonate of lime; when this crust is broken it falls to the bottom, and another, succeeds it; and in this manner the whole of the lime is soon precipitated, by absorbing carbonic acid from the air.

6. Lime is not acted on by light, neither does it combine with oxygen.

7. Sulphur and phosphorus are the only simple com- Of combosbustibles with which it unites.

Sulphuret of lime may be formed by mixing its two Sulphure. component parts, reduced to a powder, and heating them in a crucible. They undergo a commencement

by my trials, lime water contains only one 758th of its weight of

becomes red hot, raise the tube, and draw it along the coals till that part of it which contains the phosphorus is exposed to a red heat. The phosphorus is immediately volatilized, and passing through the hot lime combines with it. During the combination the mass becomes of a glowing red heat, and a quantity of phosphureted hydrogen gas is emitted, which takes fire when it comes into the air. This curious process was contrived by Dr Pearson, to whom we are indebted for the discovery of the earthy phosphurets.

Phosphuret of lime has a deep brown colour, and is moulded into the shape of the tube. It has no smell, and falls to pieces in the air. It is insoluble in water; but it has the property of decomposing that liquid. Phosphureted hydrogen gas is emitted, which takes fire as soon as it comes to the surface of the water. Part of this gas combines with the phosphuret, and forms a kind of hydrogureted phosphuret. Hence it happens, that if phospuret of lime, after being kept for some time in water, be taken out and dried, it flames when muria-

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Wan Mons has proposed the following method; but it appears to me rather inferior to that which is described in the text:—Fill a small glass matrass two-thirds with carbonate of lime in powder. Put it into a sand bath, and expose it to a heat sufficient to drive off the carbonic acid. Towards the end of the process introduce gradually a third part of phosphorus, taking care to keep the lime in a red heat. The phosphorus melts, but is prevented from burning by the remains of carbonic acid, which it disengages from the lime. When the whole of the phosphorus is introduced, shut up the matrass with a stopper, provided with a valve to let gas escape, but permiting none to enter, and let the fire be immediately withdrawn. When quite cold, the phosphuret is to be put into dry phials with ground stoppers. See Jour. de Chim. iii. 75.

tic acid is poured upon it, owing to the rapid emission of phosphureted hydrogen gas *.

8. Lime does not combine with azote; but it unites readily with muriatic acid, and forms muriate of lime.

Action of metals,

9. Lime facilitates the oxidizement of several of the metals, and it combines with several of the metallic oxides, and forms salts which have not hitherto been examined, if we except the compounds which it forms with the oxides of mercury and lead, which have been described by Berthollet.

And their exides.

The red oxide of mercury, boiled with lime-water, is partly dissolved, and the solution yields by evaporation small transparent yellow crystals. This compound has been called by some mercuriate of lime.

Lime water also dissolves the red oxide of lead, and (still better) litharge. This solution, evaporated in a retort, gives very small transparent crystals, forming prismatic colours, and not more soluble in water than lime. It is decomposed by all the alkaline sulphates, and by sulphureted hydrogen gas. The sulphuric and muriatic acids precipitate the lead. This compound blackens wool, the nails, the hair, white of eggs; but it does not affect the colour of silk, the skin, the yolk of egg, nor animal oil. It is the lead which is precipitated on these coloured substances in the state of oxide; for all acids can dissolve it. The simple mixture of lime and oxide of lead blackens these substances a proof that the salt is easily formed ‡.

10. Lime does not combine with alkalies.

Pourcrey, ii. 172. † Berthollet, Ann. de Chim. i. 62

¹ Berthollet, Ann. de Chim. i. 52.

composed of quicklime and sand reduced to a paste with water. When dry it becomes as hard as stone, and as durable; and adhering very strongly to the surfaces of the stones which it is employed to cement, the whole wall becomes in fact nothing else than one single stone. But this effect is produced very imperfectly unless the mortar be very well prepared.

The lime ought to be pure, completely free from carbonic acid, and in the state of a very fine powder: the sand should be free from clay, and partly in the state of fine sand, partly in that of gravel: the water should be pure; and if previously saturated with lime, so much the better. The best proportions, according to the experiments of Dr Higgins, are three parts of fine sand, four parts of coarser sand, one part of quick-lime recently slacked, and as little water as possible.

The stony consistence which mortar acquires is owing, partly to the absorption of carbonic acid, but principally to the combination of part of the water with the
time. This last circumstance is the reason, that if to
common mortar one fourth part of lime, reduced to
powder without being slacked, be added, the mortar
when dry acquires much greater solidity than it otherwise would do. This was first proposed by Loriot;
and a number of experiments were afterwards made by
Morveau †. The proportions which this philosopher
found to answer best are the following.

Fine sand	0.3
Cement of well baked bricks	0.3
Slacked lime	0.3
Unslacked lime	0.5

1.0

The same advantages may be attained by using as I water as possible in slacking the lime. This was pointed out by La Faye*.

Higgins found that the addition of burnt bones proved mortar by giving it tenacity, and renderin less apt to crack in drying; but they ought never exceed one fourth of the lime employed.

When a little manganese is added to mortar, it quites the important property of hardening under ter; so that it may be employed in constructing t edifices which are constantly exposed to the actic water. Limestone is found not unfrequently comb with manganese; and in that case it becomes brow calcination, instead of white. These native limest are employed for making water mortar; but good ter mortar may be made by the following process, proposed by Morveau: Mix together four parts of clay, six parts of black oxide of manganese, and 901 of limestone, all in powder. Calcine this mixture expel the carbonic acid, mix it with 60 parts of s and form it into mortar with a sufficient quantit water †.

The best mortar for resisting water is made by 1 ing with lime puzzollano, a volcanic sand brought 1

[#] Ibid. iz. 437.

[†] Ann. de Clim. xxxvii. 259.

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Italy. Morveau informs us that basakes, which is very common in this country, may be substituted for puzzollano. It must be heated in a furnace, thrown while red-hot into water, and then passed through a sieve to reduce it to the proper size ".

12. With respect to the composition of lime, nothing but Attempts conjecture was known till lately. From the theory of pose lime. Beccher and Stahl, it was the general opinion of chemists that the earths and metallic oxides are of a similar nature. Neuman, in consequence of an experiment of Henkel, attempted to obtain a metal from chalk by heating it along with combustible substances; but his experiments did not succeed †. The idea, however, was not abandoned; for it was a favourite notion of Lavoisier that all the earths might be metallic oxides 1. About the year 1790, soon after the publication of Mr Lavoisier's book, Mr Tondi and Professor Ruprecht, both of Schemnitz, announced that they had obtained from parytes, by a strong heat, a metal which they called warbonium, of the colour of iron, and attracted by the magnet; from magnesia another, which they call auswrum; a third from lime, also called austrum; and a Sourth from allumina, which they denominated apulum. Their method of proceeding was to apply a violent heat to the earths, which were surrounded with charcoal in a Hessian crucible, and covered with calcined bones in powder. But these experiments were soon after repeated by Klaproth, Savoresi, and Tihauski; and these accurate chemists soon proved that the pretended metals

[·] Ann de Chim, 222vis. 262. † Lewis, Neuman's Chemistry, p. 16. (Chemistry, p. 217. English Transf.

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were all of them phosphurets of iron. The iron, by the violence of the heat, had been extracted from the crucible, and the phosphorus from the bones.

Still more lately Desormes and Morveau were led to infer from some experiments, that lime is composed of carbon, azote, and hydrogen, and magnesia of lime and azote. But the experiments of Dorracq have demonstrated that the results obtained by these chemists were owing to the impurity of the substances on which they operated †.

It is a metallic perexide.

Mr Davy's discovery, that the fixed alkalies are metallic oxides, naturally led to the conclusion that the alkaline earths which resemble them so closely are simi-He accordingly exposed them in valar compounds. rious states to the action of a galvanic battery, and found reason to conclude that his opinion of their nature was seen correct, though he found it much more difficult to decompose them and obtain their bases, than it had become to obtain potassium and sodium by similar processes. When acted upon under naphtha they were not distinct ly decomposed. When fused with an excess of potash, and acted upon in that state, the results were rather more distinct, metallic substances appeared less fusible than potassium, which burnt the instant after they were formed, and produced a mixture of potash and the earth employed. When the earths were exposed to the action of electricity, mixed with the oxides of mercury, tin, lead, or silver, amalgams were produced, which exhibited properties indicating unequivocally, that they were mixtures of the base of the earth employed, and of the

^{*} Mem. del Instit. iii. 321.

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base of the metallicoxide mixed with it. But the quantity of amalgam obtained was too minute to admit of an accurate examination. While Mr Davy was engaged in these experiments, he received a letter from Professor Berzelius of Stockholm, informing him, that he and Dr Pontin had succeeded in decomposing barytes and lime, by negatively electrifying mercury in contact with them. Mr Davy immediately repeated this happy experiment, and succeeded completely in obtaining amalgams of the base of the earth employed.

To procure these amalgams in sufficient quantity for distilling off the mercury, and obtaining the base of the earth pure, Mr Davy combined his own previous method with that of Berzelius and Pontin. The earth was slightly moistened, mixed with one third of red oxide of mercury, and placed upon a plate of platinum connected with the positive end of the battery. A cavity was made in the earthy mixture, a globule of about 60 grains of mercury put into it, and this globule connected with the negative end of the battery by a platinum wire. The amalgams obtained in this way were distilled in glass tubes filled with the vapour of naphtha. greater part of the mercury was easily driven off, but it was extremely difficult to separate the whole. The globule that remained behind was in all cases white like silver, solid, and extremely combustible. When exposed to the air it absorbed oxygen, and regenerated the earth from which it was obtained in a few minutes.

The metallic basis of lime Mr Davy has called cal- Calcium com. He did not succeed in investigating its properties. It is white like silver, solid, and probably 4 or 8 times beavier than water. When heated in the open air it burns brilliantly, and quicklime is produced.

Neither did Mr Davy succeed in his attempts to ascertain the proportion of the constituents of lime, though he was satisfied that calcium by absorbing oxygen is converted into lime *.

SECT. II.

OF MAGNESIA.

History.

About, the beginning of the eighteenth century, 2—Roman canon exposed a white powder to sale at Rome == as a cure for all diseases. This powder he called magnesia alba. He kept the manner of preparing it a profound secret; but in 1707 Valentini informed the public that it might be obtained by calcining the lixivium which remains after the preparation of nitre †; and two years after, Slevogt discovered that it might be precipitated by potash from the mother ley ‡ of nitre §. This powder was generally supposed to be lime till Frederic Hoffman observed that it formed very different combi-

Davy's Electrockemical Researches on the decomposition of the earths, Ge. Phil. Truns. 1808.

[†] De Magnesia Alba.

The mother ley is the liquid that remains after as much as possible of any salt has been obtained from it. Common salt, for instance, is obtained by evaporating sea-water. After as much salt has been extracted from a quantity of sea-water as will crystallize, there is still a portion of liquid remaining. This portion is the mother ley.

⁵ Dist. de Magnesia Aiba.

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tions with other bodies . But little was known secraing its nature, and it was even confounded with by most chemists, till Dr Black made his celebrated eriments on it in 1755. Margraff published a disration on it in 1750 t, and Bergman another in 1775, which he collected the observations of these two losophers, and which he enriched also with many ditions of his own 1. Butini of Geneva likewise dished a valuable dissertation on it in 1779.

As magnesia has never yet been found native in a Prepare of purity, it may be prepared in the following man-Sulphate of magnesia, a salt composed of this earth sulphune acid exists in sea-water, and in many ngs, particularly in some about Epsom; from which umstance it was formerly called Eprom sale. This is to be dissolved in water, and half its weight of sh added. The magnesia is immediately precipid, because potash has a stronger affinity for sulphusacid. It is then to be washed with a sufficient quanof water, and dried.

Magnesia thus obtained is a very soft white pow- Properties. which has very little taste, and is totally destitute. mell. Its specific gravity is about 2.3 . It conto delicate vegetable blues (paper, for instance, staine with the petals of the mallow) to green.

It is not melted by the strongest heat which it has m possible to apply; but Mr Darcet observed that. wery high temperature, it became somewhat agglused. When formed into a cake with water, and

³⁰⁶s. Phys. Chim. 1722, p. 105, and 177. † Opus, ii. 90. 18 Ibrd. 1. 365. 5 Kirwan's Miner. 1. 8.

then exposed to a violent heat, the water is gradually driven off, and the magnesia contracts in its dimension at the same time, as Mr Tingry informs us, it acquires the property of shining in the dark when rubbed upon a hot iron plate.

Action of water,

3. It is almost insoluble in water: for, according to Mr Kirwan, it requires 7900 times its weight of water at the temperature of 60° to dissolve it. It is capable, however, of combining with water in a solid state, like lime; for 100 parts of magnesia, thrown into water, and then dried, are increased in weight to 118 parts. Even when combined with carbonic acid (for which it has a strong affinity) it is capable of absorbing and retaining 1; times its own weight of water without letting go a drop; but on exposure to the air, this water-evaporates, though more slowly than it would from lime.

Magnesia has never yet been obtained in a crystallized form.

When exposed to the air, it attracts carbonic acid gas and water; but exceedingly slowly. Butini left a quantity of it for two years in a porcelain cup merely covered with paper; its weight was only increased at part.

Oxygen,

4. Magnesia does not combine with oxygen, nor it altered by any of the compounds into which oxygen enters:

Simple combusti-bles,

5. The only one of the simple combustibles with which it can be united is sulphur. No person has hitherto succeeded in forming a phosphuret of magnesia.

Bergman, i. 371.

in this respect it differs from the other three alkaline Chap. III. earths.

The sulphuret of magnesia may be formed by exposing a mixture of two parts of magnesia and one part of sulphur to a gentle heat in a crucible. The result is a yellow powder, slightly agglutinated, which emits very little sulphureted hydrogen gas when thrown into water. A moderate heat is sufficient to drive off the sulphur *.

6. Magnesia does not combine with azete; but it Icombustienites with muriatic acid, and forms a compound called mariate of magnesia.

7. Magnesia has no action upon the metals; nor Metals, does it combine, as far as is known at present, with the metallic oxides, unless some intermediate substance be present.

8. Magnesia does not combine with the fixed alkalies, Alkalies. seither are its properties altered by these bodies: but it has a strong propensity to enter into triple compounds With ammonia.

9. Mr Kirwan has shown that there is but little afinity between strontian and magnesia. They do not nelt when exposed to a strong heat, at least when the Erontian exceeds or equals the magnesia +.

Equal parts of lime and magnesia, mixed together, and exposed by Lavoisier to a very violent heat, did not melt; neither did they melt when Mr Kirwan placed them in the temperature of 150° Wedgewood. The Sollowing Table, drawn up by Mr Kirwan from his own

^{*} Fourcroy, it. 165.

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experiments, shows the effect of heat on these two earths! mixed together in different proportions.

Proportions.	Heat.	Effect.
80 Lime 20 Mag.	150° Wedg.	Went through the crucible.
75 Lime 25 Mag.	160	Went through the crucible.
66 Lime 33 Mag.		Went through the crucible.
20 Lime 80 Mag.	165	Did not melt.
33 Lime 66 Mag.	138	Did not melt.
30 Lime 10 Mag.	156	Melted into a fine greenish yellow glass; but the crucible was corroded throughout.

Magnesia is used only in medicine. It is administered internally to remove acidity in the stomach.

Decompo-

Magnesia, like lime, is a metallic peroxide. Magnesia, like lime, is a metallic peroxide. Magnesia in decomposing it by the same proces by which he decomposed the other alkaline earths. When moistened magnesia is exposed to the action of galvanism in contact with mercury, the earth is reduced, and its base amalgamated with mercury much more slowly than the other alkaline earths; owing probably to its insolubility in water. The process succeeds much more rapidly when moistened sulphate of magnesia is substituted for the pure earth. To the base of magne-

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thus obtained, Mr Davy has given the name of magmon. It is a white solid metal, having the appearance silver; sinks rapidly in water, and of course is con-Merably heavier than that liquid. When the amalto of magnium is distilled in a glass tube filled with wapour of naphtha, the metal appears to act upon glass before the whole of the mercury is separated tom it. Of course it is difficult to obtain it in a state purity. When exposed to the air it rapidly absorbs gen, and is converted into magnesia. It decomses water, separating the hydrogen, and combining the oxygen; but not nearly so rapidly as the ther metals obtained from the alkaline earths; owing countless to the insolubility of the magnesia in water. but when the water is acidulated with sulphuric acid; a decomposition of water and the formation of magnee goes on with great rapidity

SECT. III.

BARTTES.

GARYTES was discovered by Scheele in 1774; and the History that account of its properties published by him in his Dissertation on Manganese f. There is a very heavy Sineral, most frequently of a flesh colour, of a foliated

Description of the Researches at the Description of the Reflies A. P. L. Fran. 1808.

^{\$} Scheele, a 61 and 28. French Translation.

texture and brittle, very common in Britain and most other countries, especially in copper mines. known by the name of ponderous spar, and was supposed to be a compound of sulphurie acid and lime. Gahn analysed this mineral in 1775, and discovered that it is composed of sulphuric acid and the new earth discovered by Scheele *. Scheele published an account of the method of obtaining this earth from ponderous spar +. The experiments of these chemists were confirmed by Bergman 1, who gave the earth the name of terms has derasa. Morveau gave it the name of barots, and Kinwan of baryers \(\); which last was approved of by Bergman #, and is now universally adopted. Different processes for obtaining barytes were published by School, Bergman, Wiegleb, and Afswelius; but little addition was made to the properties ascertained by the original discoverer, till Dr Hope published his experiments is 1793 ¶. In 1797, our knowledge of its nature was still farther extended by the experiments of Pelletier, Fourcroy, and Vauquelin **.

Prepara-

Ì

sulphate of barytes, as it is now called, by the following process, for which we are indebted to Scheele and Vanquelin. Reduce the mineral to a fine powder; mix it with the eighth part of its weight of charcoal powder, and keep it for some hours red hot in a crucible, and it will be converted into sulphuret of barytes. Dissolve the sulphuret in water, and pour nitric acid into the

[#] Bergman's Notes on Scheffer, § 167.

^{. 1742} Greife Amale, iii. 3. Eng. Trans.

¹ Opusc. iii. 191.

[§] From Bapus, heavy.

[|] Opense iv. 265.

⁹ Edin. Trens. iv. 360.

^{:44} Am. de Glim. 111 113 and 276.

pintion, and the sulphur will be precipitated. The so- Chap. 141. ution, which consists of nitric acid combined with baevers, is to be filtered and evaporated slowly till it crysdires. Put the crystals into a crucible, and expose am gradually to a strong heat; the nitric acid is drion off, and the barytes remains in a state of purity ...

Another method, attended with less expence, was pinted out long ago by Dr Hope, and afterwards imgoved by Pelletier. The method is this: Decompose sulphate of barytes by heating it strongly along ith charcoal powder. The product is to be treated th water to dissolve every thing that is soluble; and be liquid, being filtered, is to be mixed with a solution carbonate of soda. A white powder falls, Wash is powder, make it up into balls with charcoal, and eat it strongly in a crucible. When these balls are rested with boiling water, a portion of barytes is diswed, which crystallizes as the water cools.

2. Barytes obtained by the first method is a greyish- Fropertie. white, porous body, which may be very easily reduced powder. It has a harsh and more caustic taste than me; and when taken into the stomach proves a most olent poison. It has no perceptible smell. It tinges agerable blues green, and decomposes animal bodies he the fixed alkalies, though not with such energy.

Its specific gravity, according to Foureroy +, is 4; but scording to Hassenfratz only 2.374 1. But there is ason to conclude, from the method employed by this

Fquecroy, ii. 189.

1 Ann. de Chim. Expili. 2 &

When thus prepared, it always contains about 0.08 of embassis of sytes, and sometimes much more.

philosopher, that the specific gravities which he assigns are all too low.

When heated it becomes harder, and acquires internally a bluish-green shade. When exposed to the blow-pipe on a piece of charcoal, it fuses, bubbles up, and runs into globules, which quickly penetrate the charcoal. This is probably in consequence of containing water, for Lavoisier found barytes not affected by the strongest heat which he could produce.

When exposed to the air, it immediately attracts moisture; in consequence of which it swells, heat is evolved, and the barytes falls to a white powder, just as happens to quicklime when water is sprinkled on it. After the barytes is thus slacked, it gradually attracts carbonic acid, and loses its acrid properties, its weight being increased 0-22 \(\frac{1}{2}\). It cannot therefore be kept pure except in close vessels.

Action of water,

3. When a little water is poured upon barytes, it is slacked like quicklime, but more rapidly, and with the evolution of more heat. The mass becomes white, and swells considerably. If the quantity of water be sufficient to dilute it completely, the barytes crystallizes in cooling, and assumes the appearance of a stone composed of needle-form crystals; but when exposed to the air, it gradually attracts carbonic acid, and falls to powder j.

Water is capable of dissolving 0.05 parts of its weight of barytes. This solution, which is known by the name of barytes water, is limpid and colourless, has

[·] Foureroy and Vauquelia, Ann. de Chim. xxi. 276.

[†] Id. ibid. ii. 59.

¹ Mani de l'Instit. ii. 5 9.

⁶ Fourcroy, ii. 193.

Chap, III.

green, and then destroys them. When exposed to the air, its surface is soon covered with a stony crust, consisting of the barytes combined with carbonic acid.

Boiling water dissolves more than half its weight of barytes. As the solution cools, the barytes is deposited in crystals; the shape of which varies according to the rapidity with which they have been formed. When most regular, they are flat hexagonal prisms, having two broad sides, with two intervening narrow ones, and terminated at each end by a four-sided pyramid, which in some instances constitutes the larger part of the crystal. When formed slowly they are distinct and large; but when the water is saturated with barytes, they are deposited rapidly, and are generally more slender and del cate. Then, too, they are attached to one another in such a manner as to assume a beautiful foliaceous appearance, not unlike the leaf of a fern.

These crystals are transparent and colourless, and appear to be composed of about 53 parts of water and 47 of barytes. When exposed to the heat of boiling water, they undergo the watery fusion; that is to say, the water which they contain becomes sufficient to keep the barytes in solution. A stronger heat makes the water fly off. When exposed to the air, they attract carbonic acid, and crumble into dust. They are soluble in 17; parts of water at the temperature of 60°; but boiling water dissolves any quantity whatever: the reason of which is evident; at that temperature their own water of crystallization is sufficient to keep them in solution †.

" Hope, E die Tree, iv 16

+ Ibid

Book II.
Division I.
Action of exygen,
Of the simple combustibles.

- 4. Barytes undergoes no change from light; neither is it capable, as far as is known, of combining with oxygen.
- 5. None of the simple combustibles combine with it, except sulphur and phosphorus.

Sulphuret of barytes may be formed by mixing its two ingredients together and heating them in a crucible. The mixture melts at a red heat, and when cold forms a mass of a reddish-yellow colour, without any smell, which is sulphuret of barytes. This sulphuret decomposes water with great rapidity; sulphureted hydrogen is formed, which, combining with the sulphuret, converts it into a hydrogureted sulphuret. This change takes place whenever the sulphuret is moistened with water, or even exposed to the atmosphere. boiling water is poured upon sulphuret of barytes, a great quantity of sulphureted hydrogen is formed almost instantaneously, which combines with the water, and occasions the solution of the sulphuret. When the solution cools, a great number of brilliant white crystals are deposited, sometimes in the form of needles, sometimes in six-sided prisms, and sometimes in hexagonal plates. These crystals are composed of sulphureted hydrogen and barytes, and have been called by Berthollet, to whom we are indebted for the first accurate account of them, bydrosulpburet of barytes. liquid which has deposited the hydrosulphuret is of a yellow colour, and holds in solution a bydrogureted sul. phuret of barytes .

Phosphuget.

Phosphuret of barytes may be formed by putting phosphorus and barytes into a glass tube close at one

Berthollet, Ann. de Chim. xxv. 233.

ad, and heating the tube upon burning coals, as in aking phosphuret of lime. The combination takes face very rapidly. This phosphuret is of a dark brown lour, very brilliant, and very fusible. When moistenit exhales the odour of phosphureted hydrogen gas. When thrown into water, it is gradually decomposed, osphureted hydrogen gas is emitted, which takes fire then it comes to the surface of the water, and the hosphorus is gradually converted into phosphoric

Chap. 111.

6. Barytes is not acted on by azote; but it com- Action of ses readily with mariatic acid, and forms a compound ubles. led muriate of barytes.

7. Barytes has no action on metals; but it is capable combining with several of the metallic oxides, and their oxides. rming with them compounds which have not hitherto sen much examined. For instance, if it be poured into solution of silver or lead in nitric acid, it precipitates e first brown, and the second white; but if an excess barytes water be added, the precipitates are redis-Aved +.

Action of metals and

8. Barytes does not combine with the alkalies.

Of afkalies.

2. We are indebted to Bucholz for a set of experients on the action of burytes on the other earthy boles. Lime does not seem to unite with it. When ual quantities of the two earths are heated in a cruble, a mass is obtained, having some cohesion, but sopble in water 1. The phenomena were nearly the

^{🕶 🕯} aureroy, ii. 191.

Fourtroy and Vauquelin, Men. de l'Intit. ii. 61.

Buthola's Beitroge, ili. 59. He seems not to have used pure lime. tit a submuriate.

Constituents of ba-

when the resulting mass was treated with water, the barytes dissolved, but took up more of the magnesia.

10. Barytes, like the other alkaline earths, has been demonstrated by Mr Davy to be a metallic peroxide. To the metal which constitutes its base, the name of barium has been given. Mr Davy succeeded in obtaining this metal, by placing a mixture of barytes and oxide of mercury in contact with a globule of mercury, and exposing it to the action of a galvanic battery. rytes was decomposed, and its base amalgamated with the mercury. The amalgam was then heated in a glass tube filled with the vapour of naphtha, to drive off the mercury. Barium, thus obtained, is a solid metal conf the colour of silver, it melts at a temperature below redness, and is not volatilized by a heat capable of mel ing place glass, but at that temperature it acts violentl_____y upon the glass; probably decomposing the alkali of theme glass, and converting it into a protoxide. When expose d to the air it rapidly tarnishes, absorbs oxygen, and converted into barytes. It sinks rapidly in water, an seems to be at least four or five times heavier than the It decomposes water with great rapidity; hy drogen is emitted, and it is converted into barytes ---When strongly pressed it becomes flat, and hence appears to be both ductile and malleable. The proportion of oxygen necessary to convert barium into barytes has not been ascertained +.

[#] Bucholz's reitrage, iii. p. 56.

[†] Davy's Electron emical Researches on the Decomposition of the EartE.

SECT. IV.

QF STRONTIAN.

r the year 1787 a mineral was brought to Edin- History. by a dealer in fossils, from the lead mine of an, in Argyleshire, where it is found imbeded ore, mixed with several other substances. It is nes transparent and colonriess, but generally has of yellow or green. Its hardness is 5. Its speravity varies from 3.4 to 3.726. Its texture is ly fibrous; and sometimes it is found crystallislender prismatic columns of various lengths *. mineral was generally considered as a carbonate tes; but Dr Crawford having observed some difs between its solution in muriatic acid and that tes, mentioned in his treatise on muriate of baryslished in 1790, that it probably contained a new and sent a specimen to Mr Kirwan, that he might e its properties. Dr Hope made a set of expeon it in 1791, which were read to the Royal of Edinburgh in 1792, and published in the ctions about the beginning of 1794. These exnts demonstrate, that the mineral is a compound onic acid, and a peculiar earth, whose properties

· Hope, Edin. Trans. iv. 44.

are described. To this earth Dr Hope gave the name of strontites. Klaproth analysed it also in 1793, and drew the same conclusions as Dr Hope, though he was ignorant of the experiments of the latter, which remained still unpublished. Klaproth's experiments were pub-. lished in Crell's Annals for 1793 and 1794+. Kirwan also discovered the most interesting peculiarities of this new earth in 1793, as appears by his letter to Crell, though his dissertation on it, which was read to the Irish Academy in 1794, was not published till 1795. The experiments of these philosophers were repeated and confirmed in 1797 by Pelletier, Fourcroy, and Vauquelin 1, and several of the properties of the earth still farther investigated. To the earth thus detected Klaproth gave the name of strontion, from the place where it was first found; and this name is now generally adopted.

Strontian is found abundantly in different places of the world, and always combined with carbonic acid or sulphuric acid.

Prepara-

1. The carbonic acid may be expelled from the carbonate, and the strontian obtained pure, by mixing the mineral with charcoal powder, and exposing it to a heat of 140° Wedgewood §; or by dissolving the mineral in nitric acid, evaporating the solution till it crystallizes, and exposing the crystals in a crucible to a red heat till the nitric acid is driven off. Strontian may be obtained

Vol. ii. 189.

[†] Vol. i. 99. See also Klaproth's Beitroge, i. 260. and Jour. de Min. No. v. p. 61.

[?] Ann. de Chim. xxi. 113. and 276. Jour. de Min. An. vi. 3.

[&]amp; Kirwan.

from the sulphate by following exactly the process described in the last Section for obtaining barytes.

2. Strontian, thus obtained, is in porous masses, of a Properties. grevish white colour; its taste is acrid and alkaline; and it converts vegetable blues to green. Its specific gravity, according to Hassenfratz, is 1.047 . It does not act so strongly on animal bodies as barytes, nor is it poisonous +.

It does not melt when heated, like barytes; but before the blowpipe it is penetrated with light, and surrounded with a flame so white and brilliant that the eye can scarcely behold it 1.

3. When water is sprinkled on strontian it is slacked, Action of becomes hot, and falls to powder exactly like barytes; but it is not so soluble in water as that earth. One hundred and sixty-two parts of water, at the temperacure of 60°, dissolve nearly one part of strontian. The solution, known by the name of strontian water, is clear and transparent, and converts vegetable blues to a green. Hot water dissolves it in much larger quantities; and as it cools, the strontian is deposited in colourless transparent crystals. These are in the form of thin quadrangular plates, generally parallelograms, the largest of which seldom exceeds one-fourth of an inch in length. Sometimes their edges are plain, but they oftener consist of two facets, meeting together, and forming an angle like the roof of a house. These crystals generally adhere to each other in such a manner as to form a thin plate of an inch or more in length and half an inch

[·] Ann. de Chim. zzviii. 22.

⁺ Pelletier, Ibid. xxi, 150.

[!] Fourtroy, it. 227.

in breadth. Sometimes they assume a cubic form. They contain about 68 parts in 100 of water. They are soluble in 51.4 parts of water at the temperature of 60°. Boiling water dissolves nearly half its weight of them. When exposed to the air, they lose their water, attract carbonic acid, and fall into powder. Their specific gravity is 1.46 †.

Of oxygen,

- 4. Strontian is not acted on by light; neither does it combine with oxygen.
- Of combustibles
- 8. Sulphur and phosphorus are the only simple come bustibles with which it unites 1.

Sulphuret

The sulphuret of strontian may be made by fusing the two ingredients in a crucible. It is soluble in water by means of sulphureted hydrogen, which is evolved. When the solution is evaporated, bydrosulphuret of strontian is obtained in crystals, and bydrogureted sulphuret remains in solution. These three compounds resemble almost exactly the sulphuret, hydrosulphuret, and hydrogureted sulphuret of barytes; and do not therefore require a particular description. The same remark applies to the phosphuret of strontian, which may be prepared by the same process as the phosphueret of barytes §.

Phosphuret.

6. Strontian does not combine with azote; but at unites readily with muriatic acid, and forms the substance called muriate of strontian.

Action of incombustibles, metals, and alkalies.

7. Strontian has no action upon metals; but it combines with several of their oxides, and forms compounds: which have not hitherto been examined.

^{*} Hope, Edin. Trans. iv. 44. † Hassenfratz, Ann. de Chim. xxviii. E Z. † Vauquelin, Jour. de Min. An. vi. 17. § Ibid.

8. It does not combine with alkalies nor with barytes. No precipitation takes place when barytes and strontian water are mixed together *.

9. Strontian has the property of tinging flame of a beautiful red, or rather purple colour; a property discovered by Dr Ash in 1787. The experiment may be made by putting a little of the salt composed of nitric acid and strontian into the wick of a lighted candle +; or by setting fire to alcohol, holding muriate of strontian in solution. In both cases the flame is of a lively purple. In this respect it differs from barytes, which when tried in the same way is found to communicate a bluish yellow tinge to the flame 1.

Barytes and strontian resemble each other in their properties as closely as potash and soda: hence, like these two alkalies, they were for some time confounded. It is in their combination with acids that the most striking differences between these two earths are to be cobserved.

Strontian, like the other alkaline earths, has been de- Composicomposed by Mr Davy, and by a similar process. The enetallic bases of it has been called strontium. It is white, solid, much heavier than water, and bears a close resemblance to barium in its properties. When exposed to the air, or when thrown into water, it rapidly absorbs oxygen, and is converted into strontian f.

Morveau, Ann. de Chim. xxxi. 251.

Vauquelin, Jour. de Min. An. v. 10.

[?] Pelletier, Ann. de Chim. xxi, 137.

Davy's Electrochemical Researches into the Decomposition of the Bartle, We. Phil. Trees. 1808.

Back II. Division L

CHAP. IV.

OF THE EARTHS PROPER.

The Earths proper are distinguished from the alleline by neither producing any change on vegetable blues are neutralizing acids. Indeed some of them scarcely unite to these bodies. The earth's proper are five impaumber; namely,

ALUMINA,
YTTRIA,
GLUCINA,
ZIRCONIA,
SILICA.

We shall examine their properties in the following Sections.

SECT. I.

OF ALUMINA.

Alum is a salt which was well known to the ancients, and employed by them in dyeing, but they were ignorant of its component parts. The alchymists discovered that it is composed of sulphuric acid and an earth; but

be nature of this earth was long unknown. Stahl and Youman supposed it to be lime; but in 1727 Geofmay junior proved this to be a mistake, and demonstrathat the earth of alum constitutes a part of clay . 1754, Margraff showed that the basis of alum is an eth of a peculiar nature, different from every other; earth which is an essential ingredient in clays, and ves them their peculiar properties †. Hence this earth called argil; but Morveau afterwards gave it the ame of olumina, because it is obtained in the state greatest purity from alum. The properties of aluna were still farther examined by Macquer in 1758 d 1762 1, by Bergman in 1767 and 1771 |, and by scele in 1776 6; not to mention several other cheists who have contributed to the complete investigaion of this earth. A very ingenious treatise on it was ablished by Saussure junior in 1801 ¶.

Alumina may be obtained by the following process: Dissolve alum in water, and add to the solution ammona as long as any precipitate is formed. Decant off the fluid part, and wash the precipitate in a large quanty of water, and then allow it to dry. The substance has obtained is alumina; not however in a state of abolute purity, for it still retains a portion of the sulphunic acid with which it was combined in the alum. But may be rendered tolerably pure by dissolving the ewly precipitated earth in muriatic acid, evaporating the solution till a drop of it in cooling deposits small

Prepara

^{*} Men. Par 2727. † Men. Berlin, 2754 and 2759. Margraff, ii 1. † Mem. Parus. | Bergman, i. 187, and v. 71. | Scheele, i 192. French Transi. | ¶ Jours de Pôya. lii. 282.

crystals, setting it by to crystallize, separating the cry tals, concentrating the liquid a second time, and separating the crystals which are again deposited. By the process most of the alum which the earth retained we be separated in crystals. If the liquid be now mixed with ammonia as long as any precipitate appears, the precipitate, washed and dried, will be alumina near pure *.

The earth thus obtained assumes two very different appearances according to the way in which the precipitation has been conducted. If the earthy salt be dissolved in as little water as possible, the alumina has the appearance of a white earth, light, friable, very spong and attaching itself strongly to the tongue. In this state Saussure distinguishes it by the name of span alumina.

But if the salt has been dissolved in a great quantit of water, the alumina is obtained in a brittle transp rent yellow-coloured mass, splitting in pieces like n sulphur when held in the hand. Its fracture is smoo and conchoidal; it does not adhere to the tongue, a has not the common appearance of an earthy body. this state Saussure gives it the name of gelatinous and mina †.

Properties.

2. Alumina has little taste: when pure, it has smell; but if it contains oxide of iron, which it off does, it emits a peculiar smell when breathed upe known by the name of earthy smell. This smell very perceptible in common clays. The specime gravity of alumina is 2.00 ||.

A CONTRACTOR OF THE RESIDENCE OF THE PARTY O

Accum's Chemistry, ii 198 † Jour. in P. 1. 13, 100.

\$ Sevenice, Jour. de Phys. Let. 287. | Kirvini's 20, 207. 1. 1

81

3. When heat is applied to alumina it gradually loses Chan, IV. weight, in consequence of the evaporation of a quantity Contracted of water with which, in its usual state, it is combined; at the same time its bulk is considerably diminished. The spongy alumina parts with its moisture very readily, but the gelatmous retains it very strongly. Spongy alumina, when exposed to a red heat, loses 0.58 parts of its weight; gelatinous, only 0.43: Spongy alumina loses no more than 0.58 when exposed to a heat of 130° Wedgewood; gelatinous in the same temperature loses but 0.4825. Yet Saussure has shown that both species, after being dried in the temperature of 60°, contain equal proportions of water *.

Alumina undergoes a diminution of bulk proportionif to the heat to which it is exposed. This contraction seems owing, in low temperatures, to the loss of moisture: but in high temperatures it must be owing to a more intimate combination of the earthy particles with each other; for it loses no perceptible weight in any te enperature, however high, after being exposed to a heat of 130° Wedgewood t.

Mr Wedgewood took advantage of this property of Wedgealeamina, and by means of it constructed an instrument for measuring high degrees of heat. It consists of pieces of clay of a determinate size, and an apparatus for measuring their bulk with accuracy: One of these picces is put into the fire, and the temperature is estirenated by the contraction of the piece 1. The contracwion of the day-pieces is measured by means of two

[.] Tour. de Phys. 1 . 287.

⁺ Ibid.

[|] See a particular description of this thermometer in Phil. Trass. lkii. mand lerv.

Brok D. Division I. brass rules fixed upon a plate. The distance between which at one extremity is 0.5 inch, and at the other extremity 0.3 inch, and the rules are exactly 24.0 inches in length, and divided into 240 equal parts, called degrees. These degrees commence at the widest end of the scale. The first of them indicates a red heat, or 947° Fahrenheit. The clay-pieces are small cylinders, baked in a red heat, and made so as to fit P° of the scale. They are not composed of pure alumina, but of a fine white clay.

Unfortunately the contraction of these pieces is not always proportional to the degree of heat to which they have been exposed, nor do they correspond exactly with each other. The instrument, notwithstanding, is certainly valuable, and has contributed considerably towards the extension of our knowledge.

Alumina, when exposed to a very violent heat, produced by directing a stream of oxygen gas upon huming charcoal, undergoes a commencement of fusion, and is converted into a white enamel, semitransparent, and excessively hard. If we put any confidence in the calculation of Saussure, the temperature necessary for producing this effect is as high as 1575° Wedge-wood †.

Action of

4. Alumina is scarcely soluble in water, but may be diffused through that liquid with great facility. Its affinity for water, however, is very considerable. In its usual state it is combined with more than its own weight of water, and we have seen with what obstinacy

Morvenu, Jur. de l'Ecole Polytechnique, l. iii. 299. } Jour. de Phys. 2794.

it retains it. Even this combination of alumina and Chap IV. water is capable, in its usual state of drynes, of absorbing 2; times its weight of water, without suffering any to drop out. It retains this water more obstinately than any of the earths hitherto described. In a freezing cold it contracts more, and parts with more of its water, than any other earth; a circumstance which is of some importance in agriculture ".

Alumina has no effect upon vegetable blues. It cannot be crystallized artificially; but it is found native in beautiful transparent crystals, exceedingly hard, and having a specific gravity of 4. It is distinguished in this state by the name of sapphyr.

5. Alumina, as far as is known at present, is not af. Action of fected by light, neither does it combine with oxygen.

6. None of the simple combustibles are capable of Combustiuniting with it. Charcoal indeed combines with it, and bles, forms a black compound +, which is frequently found pative.

7. It is not acted on by azote; but muriatic acid lacombus dissolves it, and forms with it an uncrystallizable compound, called muriate of alumina.

8. It does not combine with metals; but it has a Metali. strong affinity for metallic oxides, especially for those exides which contain a maximum of oxygen. Some of these compounds are found native. Thus, the combination of alumina and red oxide of iron often occurs in the form of a yellow powder, which is employed as a paint, and distinguished by the name of ochre.

9. There is a strong affinity between the fixedalkalies Affaller.

Liewan's Miner. i. 9.

[†] Nicholson's Journal, ii. 101.

and alumina. When heated together, they combine and form a loose mass without any transparency. Liquid fixed alkali dissolves alumina by the assistance of heat, and retains it in solution. The alumina is precipitated again unaltered, by dropping an acid into the solution. This is the method employed by chemists to procure alumina in a state of complete purity; for alumina, unless it be dissolved in alkali, almost always retains a little oxide of iron and some acid, which disguise its properties. Liquid ammonia is also capable of dissolving a very minute proportion of newly precipitated alumina.

Barytes and strontian,

10. Barytes and strontian also combine with alumins, both when heated with it in a crucible, and when boiled with it in water. The result, in the first case, is a greenish or bluish-coloured mass, cohering but imperfectly: in the second, two compounds are formed; the first, containing an excess of alumina, remains in the state of an insoluble powder; the other, containing an excess of barytes or strontian, is held in solution by the water *.

Lime,

Alumina has a strong affinity for lime, and readily enters with it into fusion. The effect of heat on various mixtures of lime and alumina will appear from the following TABLE †.

^{*} Vanquel'n, Ann. de. Chim. xxix. 270. See also Bucholz's Beitrege, iii. 58.

[†] Kirwan, i. 65.

Proportions.	Heat.	Effect.
75 Lime 25 Alumina	150° Wedg.	Not melted.
66 Lime 33 Akumina	150	Remaineda powder.
33 Lime 66 Alumina	*	Melted.
25 Lime 75 Alumina	•	Melted.
20 Lime 80 Alumina	*	Melted.

Magnesia and alumina have no action whatever on Magnesia. each other, even when exposed to a heat of 150° Wedge-wood †.

From the experiments of Achard, it appears, that no mixture of lime, magnesia, and alumina, in which the lime predominates, is vitrifiable, except they be nearly in the proportions of three parts lime, two magnesia, one alumina; that no mixture in which magnesia predominates will melt in a heat below 166°; that mixtures in which the alumina exceeds are generally fusible, as will appear from the following Table ‡.

‡ lbid. p. 72.

^{*} These three experiments were made by Ehrman: The heat was produced by directing a stream of oxygen gas on burning charcoal, and is the most intense which it has been hitherto possible to produce.

[†] Kirwan's Miner. i, 57.

EARTES PROPER.

Book II. Division I.

3 Alumina 2 Lime 1 Magnesia	A porcelain.
3 Alumina 1 Lime 2 Magnesia	A porcelain.
3 Alumina 1 Lime 3 Magnesia	Porous porcelain.
3 Alumina 2 Lime 3 Magnesia	Porous porcelain.
3 Alumina 2 Lime 2 Magnesia	Porcelain.

None of the earths is of more importance to mankind than alumina; it forms the basis of china and stone-ware of all kinds, and of the crucibles and pots employed in all those manufactures which require a strong heat. It is absolutely necessary to the dyer and calicoprinter, and is employed too with the greatest advantage by the fuller and cleaner of cloth.

Composition of alumina. That alumina might be a metallic oxide was supported many years ago by Baron, from the resemblance between alumand the salts which sulphuric acid makes with metallic oxides. Tondi and Ruprecht announced about 1790, that they had decomposed it, and gave to the metal which they obtained from it the name of apulum. But Klaproth, Savoresi, and Tihauski demonstrated, that this pretended metal was only phosphuret of iron. Mr Davy's discovery, that the alkalies and alkaline earths

are metallic oxides, naturally led to the notion that the Chap. iV. earths proper are of the same nature. Accordingly he made a number of experiments on the subject, and has rendered the opinion probable, though he has not been able to demonstrate it in so satisfactory a manner as could have been wished. When a mixture of potash and alumina, in a state of fusion in a platinum crucible, in contact with the positive end of a galvanic battery, is touched by s platinum wire from the negative extremity, a bright combustion takes place, and metallic globules separate and burn brilliantly on the surface. The platinum. when taken out and freed from the alkali, is surrounded with metallic scales, which burn spontaneously when exposed to the air, and which decompose water with great rapidity, producing potash and an earthy matter, which was probably alumina. Potassium, amalgamated with about one third of mercury, was electrified under naphtha in contact with alumina slightly moistened. After an hour the potassium was made to decompose water, and the alkali formed neutralized. A cloudiness appeared, indicating the presence of an earthy substance. Such were the most sucessful experiments that Mr Davy was able to make in his attempts to decompose alumina and reduce it to the state of a metal. Supposing that a metallic base can be successfully extracted from it, he proposes to give that metal the name of alumium *.

Davy's Electrochemical Researches on the decomposition of the earths, IS. Pld. Trans. 1808.

SECT. II.

OF TTTRIA.

History.

Some time before 1788, Captain Arhenius discovered, in the quarry of Ytterby in Sweden, a peculiar mineral different from all those described by mineralogists. colour is greenish black, and its fracture like that of glass. It is magnetic, and generally too hard to be scratched by a knife. It is opaque, except in small pieces, when it transmits some yellow rays. Its specific gravity is 4.237 *. A description of it was published by Geyer in 1788 in Crell's Annals, and by Rinman in his Miner's Lexicon. Professor Gadolin analysed this mineral in 1794, and found it to contain a new earth: but though his analysis was published in the Stockholm Transactions for 1794, and in Crell's Annals for 1796, it was 'some time before it drew the attention of chemical mineralogists. The conclusions of Gadolin were confirmed by Ekeberg in 1797, who gave to the new earth the name of yttria+. They were still farther confirmed and extended by Vauquelin in 1800 1, and likewise by Klaproth about the same time ||; and Eke-

Gadolin, Crell's Annals, 17.6, i. 313.—Vauquelin, Ann. de Clim. xxxvi. 146.—Klaproth's Beitrage, iii. 58.

[†] Crell's Annals, 1799, ii. 63. ‡ Ann. de Chim. xxxvi. 143.

[.] I lbid. xxxvii. 86. and Beitrage, iii. 52.

berg has published a new dissertation on the subject in the Swedish Transactions for 1802 . We may therefore consider the peculiar nature of yttria as sufficiently established.

Chap. IV.

1. Hitherto yttria has been found only in the black Preparamineral first analysed by Gadolin, and hence called Gadolinite, in which it is combined with black oxide of iron and the earth called silica; and in yttrotantalite, which from the description of Ekeberg is a compound of tantalum and yttria. Both of these minerals occur only in the quarry of Ytterby. From the first, which is the most common, the earth may be procured by treating the mineral reduced to powder with a mixture of pitric and muriatic acids, till it is completely decomposed; then filtering the solution, previously evaporated nearly to dryness, and then diluting it with water. By this process the silica is left behind. The liquid which passes through the filter is to be evaporated to dryness. and the residue heated to redness for a considerable time in a close vessel, and then redissolved in water and filtered. What passes through the filter is colourless; when treated with ammonia, pure yttrea falls.

2. Yttria thus procured has the appearance of a fine Properties. white powder, and has neither taste nor smell. It is not melted by the application of heat. It has no action on regetable blues. It is much heavier than any of the other earths, its specific gravity, according to Ekeberg, being no less than 4.842.

It is insoluble in water; yet it is capable of retaining a great proportion of that liquid, as is the case with

⁹ Jour de Chim. iii 78.

alumina. Klaproth ascertained, that 100 parts of yttria, precipitated from muriatic acid by ammonia, and dried in a low temperature, lose 31 parts, or almost a third of their weight, when heated to redness in a crucible. Now this loss consists of pure water alone.

It is not soluble in pure alkalies; but it dissolves readily in carbonate of ammonia, and in all the other alkaline carbonates. It combines with acids, and forms with them salts which have a sweet taste, and at the same time a certain degree of austerity. Some of these salts have a red colour. Yttria is the only earthy body known which has the property of forming coloured salts with acids.

¥

3. Yttria is not altered by light, nor is it likely that it combines with oxygen. From the experiments of Klaproth, it does not appear to combine readily with sulphur; nor is it likely that it unites with any of the other simple combustibles.

We may take it for granted that it is not affected by azote; but it combines with muriatic acid, and forms a salt not capable of crystallizing. Its action on the metals and metallic oxides is unknown.

Composi-

No attempts have been made to decompose yttria.

But from analogy there can be little doubt that, like the sother earths, it is in reality a metallic peroxide. The experiments of Ekeberg, if correct, may be considered as a demonstration of the truth of this opinion. When he treated yttria with muriatic acid, oxymuriatic acid is said to have been evolved.

Chap. IV.

SECT. III.

GLUCINA.

beryl is a transparent stone, of a green colour, History. considerable degree of hardness, which is found malized in the mountains of Siberia, and in many parts. Vauquelin analysed this mineral in 1798, e request of Hauy, to determine whether it was ed of the same ingredients with the emerald, as y had conjectured from mineralogical consider-The result of the analysis was a confirmation suspicions of Hany, and the discovery of a new to which Vanquelin and his associates gave the of glucina. The experiments of Vauquelin have repeated by Klaproth † and other eminent che-

To obtain glucina pure, the beryl or the emerald, Preparaced to powder, is to be fused with thrice its weight tash. The mass is to be diluted with water, dised in muriatic acid, and the solution evaporated to bess. The residuum is to be mixed with a great mity of water, and the whole thrown on a filter. silica, which constitutes more than half the weight the stone, remains behind; but the glucina and the earths, being combined with muriatic acid, remain

in solution. Precipitate them by means of carbonate of potash. Wash the precipitate, and then dissolve it in sulphuric acid. Add to the solution sulphate of potash; evaporate it to the proper consistency, and set it by to crystallize. Alum crystals gradually form. When as many of these as possible have been obtained, pour into the liquid carbonate of ammonia in excess, then filter, and boil the liquid for some time. A white powder gradually appears, which is glucina.

Properties.

2. Glucina, thus obtained, is a soft light white powder, without either taste or smell; which has the property of adhering strongly to the tongue. It has no action on vegetable colours. It is altogether infusible by heat; neither does it harden or contract in its dimensions, as is the case with alumina. Its specific gravity is 2.976 *.

It is insoluble in water, but forms with a small quantity of that liquid a paste which has a certain degree of ductility.

- 3. It does not combine with oxygen nor with any of the simple combustibles; but sulphureted hydrogen dissolves it, and forms with it a hydrosulphuret, similar to other hydrosulphurets in its properties †.
- 4. Azote has no action on it; but muriatic acid dissolves it, and forms with it a sweet-tasted salt, called muriate of glucina.
- 5. Glucina is soluble in the liquid fixed alkalies, in which it agrees with alumina. It is insoluble in ammonia, but soluble in carbonate of ammonia, in which

^{*} Ekeberg, Ann. de Clim. > liii. 277. † Fourcrey, ii. 159.

respect it agrees with yttria; but it is about five times more soluble in carbonate of ammonia than that earth.

It combines with all the acids, and forms with them aweet-tasted salts", as is the case also with yttria.

Mr Davy tried to decompose glucina by the same Composimeans as have been related in the first section of this Chapter with respect to alumina. The effect was the same. We have reason therefore to believe that it is a metallic peroxide. Mr Davy proposes to call its base glucium.

SECT. IV.

OF ZIRCONIA.

Anong the precious stones which come from the island History. Ceylon, there is one called jargon or mircon, which is mossessed of the following properties.

Its colour is various; grey, greenish-white, yellowash, reddish-brown, and violet. It is often crystallized, either in right angular quadrangular prisms surmounted with pyramids, or octahedrons consisting of double quadrangular pyramids. It has generally a good deal of lustre, at least internally. It is mostly semitransparent. Its hardness is from 10 to 16. Its specific gravity from 4.416 to 4.7 †.

It loses scarcely any of its weight in a melting heat;

[·] Hence the name glucing, from yourse, exceet

[†] Kiewan's Miner. 1. 333.

3° seci "acurer"

or silectors, who inalysed it is 1789, found that 300 grams, ther emanant wit for an hour and a half, was me me lourth of a grain igner than at that . No. her was t stacked wher he mariate or sulphair and, even when issisted by bear. At last, by calcining t with a wife mantity of come be dissolved is in there will and bound that \$100 meets of it contained says of olica. 107 st a mixture of melant and iron, and 60 of r very meth, museuset of reculiar momentus; which has received he same it sarrows, from the mineral is which t was intected. County probably to the seasons of the aircon, aonone attenument to remain the maintain of Klaaroth, ar a versty his discovery. In 1795 he publish ed his mairas it the inaccess, monther mineral from the same island, in which he also interest a large meportion of arconia, expressing his larges that it would induce chemists to turn their attenuous to the subject ! This molest induced Covers Moreover, in 1798, tour mine the avacuates of Experity in France. Therepas red similar to the systematic of Ception, and contained the geograph of sirranus inclusion by Commentit. Frage experiments were more with represent, with congre of the new earth still institut communic be Touggelin !.

Fragme

Comming the historie been found only in the sine and and the highest with the highest process of process, while the highest the mineral to proceed, with the or we give of process, and fore it is a control.

Only the or was give of process, and fore it is a control.

Only the or was give of process, and fore it is a control.

⁴ Jour de Phys. RERVI. 180

[†] Beitrage, i 23

¹ Ann. de (48m. 22i. 72.

[§] Ibid. axis 158, and Jour. de Min At. v 97.

potash is extracted; then dissolve the residuum as possible in diluted muriatic acid. Boil the soluto precipitate any silica which may have been dised; then filter, and add a quantity of potash. The mia precipitates in the state of a fine powder.

Zirconia, thus prepared, has the form of a fine properties. powder, which feels somewhat harsh when rubbetween the fingers. It has neither taste nor odour. infusible before the blowpipe; but when heated ently in a charcoal crucible, it undergoes a kind of exfect fusion, acquires a grey colour, and something e appearance of porcelain. In this state it is very d. its specific gravity is 4.3, and it is no longer soe in acids.

sirconia is insoluble in water; but it has a consideraffinity for that liquid. When dried slowly, after precipitated from a solution, it retains about the of its weight of water, and assumes a yellow coand a certain degree of transparency, which gives great resemblance to gum arabic .

It does not combine with oxygen, simple combusazote, nor metals; but it has a strong affinity for metallic oxides, especially for oxide of iron, which it is very difficult to separate it.

is insoluble in liquid alkalies, neither can it be fusolong with them by means of heat; but it is soluble Pealine carbonates.

Scarcely any experiments have been made to ascerits affinity for the different earths. It is known, ever, that a mixture of alumina and zirconia is cae of fusion.

[·] Vauquelin, Ann. de Ch m. 1x1 158

Zirconia combines with all the acids, and forms salts; which have a peculiar astringent taste, and are many of them insoluble in water.

This earth has not hitherto been applied to any use. Its scarcity, and the difficulty of procuring it in a state of purity, exclude it at present from any chance of being employed for the purposes of domestic economy.

Composi-

The same attempts were made by Mr Davy to decompose zirconia as those made upon alumina and glucina, and with similar results. The metallic base he proposes to call zirconium.

SECT. V.

OF SILICA.

History.

There is a very hard white stone, known by the name of quarts, very common in almost every part of the world. Sometimes it is transparent and crystallized, and then is called rock crystal. Very frequently it is in the form of sand. As this stone, and several others which resemble it, as flint, agate, calcedony, &c. have the property of melting into a glass when heated along with fixed alkali, they were classed together by mineralogists under the name of vitrifiable stones. Mr Pott, who first described their properties in 1746, gave them the name of siliceous stones, on the supposition that they were all chiefly composed of a peculiar earth called siliceous earth or silica. This earth was known to Glau-

Chap. IV.

escribes the method of obtaining it but it fore its properties were accurately ascertainroy * endeavoured to prove that it thight be nto lime, and Pott + and Baumé I that it proverted into alumina: but these assertions i by Cartheuser §, Scheele !!, and Bergman ¶. chemist we are indebted for the first accuof the properties of silica **.

may be obtained pure by the following pro- Preparatogether, in a crucible, one part of pounded rtz, and three parts of potash, and apply & ent to melt the mixture completely. Dismass formed in water, saturate the potash tic acid, and evaporate to dryness. Towards the evaporation the liquid assumes the form and when all the moisture is evaporated, a remains behind. This mass is to be washe quantity of water and dried; it is then site of purity.

, thus obtained, is a fine white powder, with. Properties aste or smell. Its particles have a harsh feel, consisted of very minute grains of sand. Its vity is 2.66 ††.

e subjected to a very violent heat without unny change. Lavoisier and Morveau exposed action of a fire maintained by oxygen gas y alteration 11. Saussure indeed has suc-

Par. 1746, p. 286. + Litbogn. p. 3. Praf. Miner Abb. de Cbym. c, i. 191. s Terres Geoponiques, 1773, Opuse. v. 59. †† Kirwan's Min. i. 10. ıc. ii. 26 de l' Ecole Polytecha. I. iii. 299. Z.

Book II. Division I.

of it so extremely minute as scarcely to be perceptible without a glass. According to the calculation of this philosopher, the temperature necessary for producing this effect is equal to 4043° Wedgewood.

Action of water.

3. It is insoluble in water except when newly precipitated, and then one part of it is soluble in 1000 parts of water *. It has no effect on vegetable colours.

It is capable of absorbing about one-fourth of its weight of water, without letting any drop from it; but on exposure to the air, the water evaporates very readily †. When precipitated from potash by means of muriatic acid and slow evaporation, it retains a considerable portion of water, and forms with it a transparent jelly; but the moisture gradually evaporates on exposure to the air.

Silica may be formed into a paste with a small quantity of water; this paste has not the smallest ductility, and when dried forms a loose, friable, and incoherent mass ‡.

Silica is capable of assuming a crystalline form. Crystals of it are found in many parts of the world. They are known by the name of rock crystal. When pure they are transparent and colourless like glass: they assume various forms; the most usual is a hexagonal prism, surmounted with hexagonal pyramids on one or both ends, the angles of the prism corresponding with those of the pyramids. Their hardnes is very great, amounting to 11. Their specific gravity is 2.653 ||.

^{*} Kirwan's Min. i. 10.

[†] Ibid.

¹ Scheele.

[#] Kirwan's-Mis. i. 242.

Chap. IV.

The first method was discovered by Bergman. solved silica in fluoric acid, and allowed the solutemain undisturbed for two years. A number talls were then found at the bottom of the vessel, of irregular figures, but some of them cubes tair angles truncated. They were hard, but not ompared in this respect with rock crystal. other method was discovered by accident. Proseigling of Erfurt had prepared a liquor silicum, was more than usually diluted with water, and ed a superabundance of alkali. It lay undistor eight years in a glass vessel, the mouth of

for eight years in a glass vessel, the mouth of was only covered with paper. Happening to it by accident, he observed it to contain a num-crystals; on which he sent it to Mr Tromms-professor of chemistry at Erfurt, who examined le liquor remaining amounted to about two ounces. Take was covered by a transparent crust, so that the vessel might be inverted without spilly of the liquid. At the bottom of the vessel were her of crystals, which proved on examination to phate of potash and carbonate of potash. The mathetop consisted partly of carbonate of potash, of crystallized silica. These last crystals had asthe form of tetrahedral pyramids in groupes; were perfectly transparent, and so hard that they

Bilica neither combines with oxygen, with the

Action of simple substances,

fire with steel 1.

Bergman, ii. 32.

Potash combined with sulphuric acid and carbonic acid

[!] Nicholson's Journal, 1. 227.

Book II. Division I. with many of the metallic oxides by fusion, and forms various coloured glasses and enamels.

5. Azote has no action on silica; neither has muriatic acid when the silica is in a solid state; but when the silica is combined with an excess of alkali, muriatic acid dissolves the compound, and forms a permanent solution. By concentrating this solution, the silica separates from it in the form of a jelly.

Action of

o. There is a strong affinity between silica and fixed alkalies. It may be combined with them either by fasing them along with it in a crucible, or by boiling the liquid alkalies over it. When the potash exceeds the silica considerably, the compound is soluble in water, and constitutes what was formerly called liquor silicant and now sometimes silicated potash or soda. When the silica exceeds, the compound is transparent and colourless like rock crystal, and is neither acted on by water, air, nor (excepting one) by acids. This is the substance so well known under the name of glass.

Silica is not acted on by ammonia, whether in the gaseous or liquid state.

Of barytte.

7. There is a strong affinity between barytes and silica. When barytes water is poured into a solution of silica in potash, a precipitate appears, which is considered by Morveau as the two earths in a state of combination. Barytes and silica may be combined by means of heat. The compound is of a greenish colour, and soheres but imperfectly †. The effect of heat on va-

^{*} Morveau, Ann. de Chim. xxxi. 250.

f Vauguslin, ibid. xxix. 271.

BILICA.

iixtures of barytes and silica will appear from Chap. IV. owing experiments of Mr Kirwan *.

rtions.	Heat.	F.ffect.
ica rytes	150° Wedg.	A white brittle mass.
ica rytes	150	A brittle hard mass, semi- transparent at the edges.
ics rytes	150	Melted into a hard, somewhat porous porcelain
ica rytes	148	A hard mass not melted:
ica rytes	148	The edges were melted into a pale greenish matter between a porcelain and enamel.
ica rytes	150	Melted into a somewhat porous porcelain mass.
ica rytes	150	Melted into a yellowish and partly greenish white porous porcelain.

itian and silica combine with each other nearly Strontian, ame manner.

e is also an affinity between silica and lime. Lime, lime-water is poured into a solution of silica in a precipitate appears, as Stucke discovered. This ate is a compound of silica and lime +. These

van's Miner. i. 57.

din, Ann. de Chim. axii. 110.—Morveau, ibic'. axxi. 250.

Book 7. Division L

two earths may be combined also by means of heat. They form a glass, provided the quantity of lime be not inferior to that of silica. The effect of heat upon these earths, mixed in various proportions, will appear from the following experiments of Mr Kirwan.

Proportions Heat		Effect.		
50 Lime 50 Silicz	150° Wedg.	Melted into a mass of a white colour, semitransparent at the edges, and striking fire, though feebly, with steel! it was somewhat between porcelain and enamel.		
80 Lime 20 Silica	156	A yellowish white loose powder.		
20 Lime 80 Silica	156	Not melted, formed a brit- tle mass.		

Magnesia,

Equal parts of magnesia and silica melt with great difficulty into a white enamel when exposed to the most violent heat which can be produced †. They are infasible in inferior heats in whatever proportion they are mixed ‡.

And alumi-

There is a strong affinity between alumina and silica. When equal portions of silicated and aluminated potash are mixed together, a brown zone immediately appears, which may be made, by agitation, to puss through the whole liquid. After standing about an hour, the mixture assumes the consistence of jelly #.

[#] Kirwan's Min. i. 56. † Lavoisier, Men. Per. 1787, p. 598.

¹ Achard, Mem. Best. 1780, p. 33.

⁴ Morveau, Ann. de C'im. xxxi. 249.

Chap. IV.

ned into a paste with water, and dried, they d contract a considerable degree of hardness. ted in the temperature of 160° Wedgewood, me very hard, but do not fuse. Achard a infusible in all proportions in a heat proe inferior to 150° Wedgewood. But when a very strong heat, they are converted infopaque glass, or rather enamel. Porcelain, brick, tiles, and other similar substances, are chiefly of this compound. Mixtures of silinina in various proportions constitute clays; re seldom uncontaminated with some other

ws from the experiments of Achard, that of lime, magnesia, and silica, may be melt-reenish coloured glass, hard enough to strike teel; that when the magnesia exceeds either r two, the mixture will not melt; that when xceeds, the mixture seldom melts, only innim in the following proportions; three siline, one magnesia, which formed a porcelain; then the lime exceeds, the mixture is genele †.

re of silica and alumina may also be combined es or strontian by means of heat. The mix-readily into a greenish coloured porcelain ‡. e experiments of Achard and Kirwan, we in mixtures of lime, silica, and alumina, ime exceeds, the mixture is generally fusible

Min. i. 58.

^{. 1780,} p. 33. and Jour. de Plys. xxiv.

Book II. Division L either into a glass or a porcelain, according to the proportions. The only infusible proportions were,

2 | 3 ; Lime 1 | 1 | Silica 2 | 2 | Alumina.

That if the silica exceeds, the mixture is frequently fusible into an enamel or porcelain, and perhaps a glass; and that when the alumina exceeds, a porcelain may often be attained, but not a glass.

As to mixtures of magnesia, silica, and alumins, when the magnesia exceeds, no fusion takes place at 150°. When the silica exceeds, a porculain may often be attained; and three parts silica, two magnesia, and one alumina form a glass. When the alumina exceeds, nothing more than a porcelain can be produced †.

Achard found that equal parts of lime, magnesia, silica, and alumina, melted into a glass. They fused also in various other proportions, especially when the silica predominated.

Silica differs from all the other earths in not combining with any of the acids except the fluoric, phosphoric, and boracic; to which perhaps we may add the mutiatic.

Silica is one of the most important of the earths. It is the chief ingredient of those stones which seem to constitute the basis of this terrestrial globe. It is an essential ingredient in mortar, in all kinds of stoneware, and in glass.

Compositien. Mr Davy made many experiments to ascertain the composition of silica, and has rendered it probable that it is a metallic peroxide. As it approaches in some of

Kirwan's Mis. i. 73.

quisite, in the first place, to inquire whether it might not be a compound of an unknown scid and earth. But when exposed to the action of a powerful galvanic battery it was not decomposed; of course there was no reason for considering it as propable that it was of a saline nature. On the supposition that it was a metallic peroxide, it was exposed to experiments precisely similar to those formerly described, when treating of the decomposition of alumina, and the results were exactly the same. Hence there is reason to consider it as a compound of oxygen and a peculiar metal, though the experiments cannot be considered quite decisive. To the supposed metal Mr Davy proposes to give the name of efficients.

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Book II. Division L

CHAP. V.

rks on the alkalies and

Eartis_

THE terms alkali and earth were brought into me

meaning of

Original

elkeli and

earth.

long before chemistry became an accurate science. We need not therefore be surprized that their meaning was somewhat vague; and that as chemical substances multiplied, it became difficult to ascertain to which of the two they ought to be assigned. Two properties appear to have been considered as essential to the alkalies. 1. They were capable of combining with acids, and of depriving them of their acidity, while they themselves lost their characteristic properties, or were neutralized. 2. They were soluble in water. Insolubility in water, or aridity, as the chemists termed it, joined to incombustibility, was considered as characteristic of an earth. These properties served to arrange newly discovered substances. If they combined with acids, and were soluble in water, they were considered as alkalies; but if they were insoluble in water, they took their station among the earths. Hence soda, as soon as discovered, was denominated an alkali, while lime, barytes, stronwan, &cc. were successively placed among the earths; for the solubility of the first of these bodies in water was long misunderstood. The whole of the lime was not considered as soluble in water, but a certain portion of it which approached alkalies in its nature; and though his error had been rectified before the discovery of strontian, yet analogy led chemists to class this body slong with lime and barytes, which had already taken their place among earths.

The alkalies naturally subdivided themselves into sions of fixed and volutile; a distinction made at an early pe- them. siod, sufficiently correct, and attended with many ade, vantages. The subdivision of the earths into alkaline and proper was a much later improvement. Indeed it was only after the earths had been examined in a state of tolerable purity that such a subdivision was possible. Those earths that possessed all the characteristics of alkalies, excepting that of being very soluble in water. were denominated alkaline, while those that were incapable of neutralizing acids, and which were insoluble: in water, were considered as more peculiarly entitled to the name of earth. This subdivision was obviously. intended to correct the too great vagueness of the termswith and alkali as originally applied. It appears more. proper than some other methods that have been prace. ised for the same purpose. Fourcroy, for instance, . lasses barytes and strontian with the alkalies. No ault can be found with this arrangement, because the division of these bodies into earths and alkalies is perfeetly arbitrary. But surely if barytes and strontian be placed among alkalies, lime ought not to be excluded; for barytes and strontian do not possess a single alka, line property of which lime is destitute. And if lime

Book II. Division I. be reckoned among the alkalies, no good reason can be given why magnesia should be excluded. The truth is, that these bodies graduate into each other so nicely that they can scarcely be placed in different classes. This is a sufficient reason for preferring the common division to the new one proposed by Fourcroy.

Bertholiet's definition of alkali.

Berthollet has suggested another definition of alkalies, which appears to me much more correct than that of According to him, all bodies capable of Fourcroy. estralizing acids are entitled to the name of alkalies. Attalisity is the property of neutralizing acids, and acidity that of neutralizing alkalies. According to this definition, the alkalies, both fixed and volatile, and the aikaline earths, ought to be considered as alkalies. Though objections might perhaps be started to this definition, as is the case indeed with all definitions whatever, yet it must be allowed to be much more precise, and gives us a mure estisfactory notion of elkalinity than any that preceded it. But if the capacity of newtralizing acids constitutes alkalinity, it is obvious that those budies inust possess most of that property, which assissor capable of mentralizing acids, or which are capable of neutralizing the greatest quantity of acid. Accosding to this docume the alkalies arrange themselves in the following order; that one standing first which neutralizes the greatest quantity of acid *;

Ammonia,

"Magnesia,

"Time,

"Time,

"Time,

Berthollet, Statique Chemique, i. p. 125. The order was established from the experiments of Kirwan.

Soda, Potash, Strontian, Barytes.

Chap. V.

The earths proper combine with acids, but do not tralize them *.

The alkalies (using the word in Berthollet's sense) not combine with oxygen. They do not act upon lrogen nor carbon; but they combine with sulphur, unite with phosphorus, or at least act upon it with siderable energy. Their action upon metals is not y remarkable. Some metallic oxides they dissolve, le upon others they have no effect. They are all re or less soluble in water except magnesia; but the aline earths much less so than the others.

The earths proper do not unite with oxygen, at least ar as has been ascertained in a satisfactory manner. by do not unite with simple combustibles nor metals, have a much less tendency to enter into combina-

have a much less tendency to enter into combinais than the alkalies.

The following Table exhibits some of the most tarkable characters of the alkaline and proper ths †.

To this there are some exceptions. This constitutes one of the deof Berthollet's definition.

In this Table several properties have been placed, with which the er will become acquainted by perusing the subsequent part of this k. It was thought worth while to add them, that the Table might ain the most important properties of the earths collected together.

Book II. Division L

Zartha	Solubility in r of	Tinge vegetable	Soluble in potash.	Soluble in carbonate of ditte.	Soluble in carbonate of animomia.	Soluble in marianc	Procipitated by prushate of potash.	Precipitated by	Precipit by by-	Precipitated by sac-
Burytes	0-050	i				1		1		1
Stroot.	0.002	2	:			9		7		2
Lime	0-002	9				3				3
Magnes.	0.000	4				4	·			
Alum.	0.000		1			5			1	
Yttria	0.000			1	1	đ	1	1		
Gluoina	0.000		, 2	2	2	7		2		4
Zircon.	0.000			9		8	2	3	2	
Silica	0-000		9							

It deserves attention, that a considerable number of these bodies may be divided into pairs, which have a striking resemblance to each other. These pairs are,

1. { Potash Soda S. } Yttria Glucina Glucina Alumina Zirconia

But the resemblance between alumina and zirconia, which constitute the last pair, is not so close as that

After being precipitated.

between the bodies which form the other pairs. Ammonia, magnesia, and silica, have none of them corresponding substances.

Chap. V.

Now that the secret of the composition of the alkalies and earths has been revealed, it is obvious that they all belong to the same class of substances, and that they ought to be placed among the metallic oxides. Thus the conjecture of the old chemists has been verified with respect to the earths; while the constitution of the alkalies has turned out different from every previous conjecture.

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Book II. Division II.

DIVISION II.

OF

PRIMARY COMPOUNDS.

If the science of chemistry had made sufficient progress, this division would comprehend all the compounds which the simple confinable bodies are capable of forming with each other. But we are not yet sufficiently acquainted with all of these combinations to be able to detail them at length; and many substances known to exist, and to belong to this division, have not been sufficiently examined to enable us to assign them their proper situation. We cannot, therefore, introduce all primary compounds into this division, but those only, in the analysis of which chemistry has made considerable progress.

The confinable bodies were divided into four classes; namely, Oxygen, Combustibles, Incombustibles, and Metals. Oxygen unites with all the bodies belonging to the other three classes; and with most of them it unites in various proportions. United in one proportion, it forms a set of bodies incapable of neutralizing alkalies, to which the name of oxide has been assigned; united in another proportion, it forms a set of

Each of these sets comprehends under it a variety of bodies of great importance in chemistry, and which, therefore, deserve to be examined with attention. The combustibles unite with each other, and form a very numerous class of bodies still capable of combustion, and which, therefore, may be called compound combustibles. They unite also with incombustibles and with metals. The incombustibles, as far as we know, do not unite with each other, nor with the metals; but the metals combine together, and form compounds called alloys. Hence we see that the primary compounds naturally divide themselves into the six following classes:

Book II. Division II.

Division

1. Oxides;

2. Acids,

3. Compound combustibles;

- 4. Combinations of combustibles and incombustibles;
- 5. Combinations of combustibles and metals;
- 6. Combinations of metals with each other.

Of these classes, the two last, in the present state of the cience, are of least importance. The compounds belonging to them are scarcely entitled to a separate examination, and for that reason were introduced in the first Book of this Work, during the examination of the simple substances, of which they are compounds. As to the fourth class, we are at present acquainted only with one substance which we know to belong to it. That substance is ammonia, which was arranged with the fixed alkalies and earths, for reasons formerly specified. The first three classes of primary compounds alone remain to be examined. They shall form the subject of the following chapters.

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Book II. Division II.

CHAP. I.

OF OXIDES.

Combination of oxygen.

WE have seen already that oxygen combines with bodies in various proportions, constituting a variety of compounds with almost every substance with which it is capable of uniting. Now the whole of the compounds into which oxygen enters may be divided into two sets: 1. Those which possess the properties of acids; and, 2. Those which are destitute of these properties. The first set of compounds are distinguished by the term & cids; to the second, the term oxide has been appropriated. By oxide, then, is meant a substance composed of oxygen and some other body, and destitute of the properties which belong to acids. It is by no means uncommon to find a compound of the same base and oxygen belonging to both of these sets, according to the proportion of oxygen which enters into the compound. In all these cases, the smaller proportion of oxygen constitutes the oxide; the larger the acid. Hence it folthat oxides always contain less oxygen than acids the same base.

Chap. I.

e oxides, then, which we have to examine in this er, are combinations of oxygen with the simple ustibles, the simple incombustibles, and the mesubstances which may be denominated the bases: oxide. Now the oxides differ considerably from other, according to the nature of the base and the er in which the combination has been formed: of them are products of combustion, others of are combustible, while a third set are supporters of sustion. The metallic oxides, which have been already examined, belong partly to the first and partly to ast of these classes. The following Table exhibits a of the different oxides (omitting the metallic) ared under their respective classes.

I. OXIDE PRODUCTS.

Base. Name. Hydrogen Water.

II. Combustible Oxides.

Carbon......Carbonic oxide.

Phosphorus......Oxides of phosphorus.

Sulphur......Oxides of sulphur.

III. Oxide Supporters.

Azote..... SNitrous oxide.
Nitric oxide.
Muriatic acid.....Oxymuriatic acid.

examination of these oxides shall occupy our attenin the following Sections. Book II. Division II.

I. OXIDE PRODUCTS.

Hydrogen differs from the other simple combustibles in several respects. It is the only one of them which is capable of uniting with only a single dose of oxygen, and of forming a compound entirely destitute of acid properties. This compound is Water. It is formed by combustion, and is therefore a product. All the other products into which the simple combustibles enter are acids. The only oxide product, then, at present known (exclusive of the metallic) is water.

SECT. I.

OF WATER.

This well-known liquid is found in abundance in every part of the world, and is absolutely necessary for the existence of animals and vegetables. When pure, in which state it can be obtained only by distillation, it is transparent, and destitute of colour, taste, and smell.

Weight.

1. As this liquid, from the ease with which it may be procured in a state of purity, has been chosen for a standard by which the comparative weight of all other bodies may be estimated, it becomes of the greatest importance to ascertain its weight with precision: But its density varies with the temperature. At the temperature of 36°, its density is a maximum; and for the two or three degrees on each side of 36°, the varia-

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tion of its density is scarcely perceptible. Now, from the experiments of Lefevre Gineau, performed with great care, in order to ascertain the weight of the millegramme, it follows, that, at the temperature of 40°, a French cubic foot of distilled water weighs 70lbs. and 223 grains French = 529452.9492 grains troy *. Therefore an English cubic toot, at the same temperature, weighs 437102.4946 grains troy, or 999.0914161 ounces avoirdupois. Hence a cubic inch of water at 40° weighs 252.953 grains, and at 60°, 252.72 grains. A cubic foot of water, at the temperature of 55°, weighs, according to the experiments of Professor Robison of Edinburgh, 998.74 avoirdupois ounces, of 437.5 grains troy each, or only 1.26 ounces less than 1000 avoirdupois ounces: so that rain water, at the same temperature, will weigh pretty nearly 1000 ounces. cific gravity of water is always supposed = 1.000, and it is made the measure of the specific gravity of every other body.

2. When water is cooled down to 32°, it assumes Ica the form of ice. If this process goes on very slowly, the ice assumes the form of crystalline needles, crossing each other at angles either of 60° or 120°, as Mr de Mairan has remarked; and it has been often observed in large crystals of determinate figures †. Ice, while kept at a temperature considerably below 32°, is very hard, and may be pounded into the finest dust. It is elastic. Its specific gravity is less than that of water.

3. When water is heated to the temperature of 212°, Steme

Chap. I.

^{*} Jour. de Phys. xlix. 171.

[†] See observations on this subject by Grew. Phil. Trans. Abr. ii. 54.

Book II. Division II. it boils, and is gradually converted into steam. Steam is an invisible fluid like air, but of a less specific gravity. It occupies about 1800 times the space that water does. Its elasticity is so great, that it produces the most violent explosions when confined. It is upon this principle that the steam-engine has been constructed.

The phenomena of boiling are owing entirely to the rapid formation of steam at the bottom of the vessel. The boiling point of water varies according to the pressure of the atmosphere. In a vacuum water boils at 70°; and when water is confined in Papin's digester, it may be almost heated red hot without boiling. The mixture of various salts with water affect its boiling point considerably. Mr Achard made a number of experiments on that subject; the result of which may be seen in the following TABLES*.

Boiling point affected by

CLASS I. Salts which do not affect the Boiling Point.

Sulphate of copper.

CLASS II. Salts which raise the Boiling Point

ò	Muriate of soda]	10.350
d s	Sulphate of soda	the just	5.6
ate	Sulphate of potash	P G	0.0
tio	Nitrate of potash	ing	3.2
sat Lu	Boracic acid	P. E.	2.2
4	Carbonate of soda	ا هر ا	2.35

This augmentation varies with the quantity of salt

^{*} Trans. Berlin. 1785.

lved. In general, it is the greater the nearer the ion approaches to saturation.

LASS III. Salts which lower the Boiling Point.

(In a small quantity, lowers the boil-	
In a small quantity, lowers the boil- ing point	50
(Saturated solution of	2
hate of magnesia, Saturated solution of 2.4 Saturated solution of 1.1	7
A greater quantity of 0.07 A saturated solution of 0.0	!
hate of lime, bhate of zinc, hate of iron, tate of lead, in any proportion, 10.02 10.24 10.24	2 5 2

CLASS IV.

riate of monia,	Small quantity of, lowers the boiling point0.45° Saturated solution of, raises do.9.79
bonate of otash,	Small quantity of, lowers do. 0.45 Saturated solution of, raises do. 11.2

Vater was once supposed to be incompressible; but contrary has been demonstrated by Mr Canton. : Abbé Mongez made a number of experiments, gafter that philosopher, on the same subject, and uned similar results.

. Water is not altered by being made to pass through Action of d hot tube. Heat does not seem capable of decom ing it; neither is it affected by the action of light.

. It has the property of absorbing atmospheric air; Absorbe it always contains a portion of it when it has been

Book II. Division IL exposed to the atmosphere. The greater part of this air is driven off by boiling: but, from the experiments of Dr Priestley, it appears that the whole of it is not separated; nor can it be completely separated without great difficulty. Water owes its agreeable taste to the presence of air; hence the insipidity of boiled water, It absorbs oxygen gas in preference to air, and nearly in the same proportion, as was first ascertained by Scheele.

How air may be separated rom it.

Mr Driessen has shown, that in order to free water from air, it must be boiled at least for two hours, and kept in a flask with its mouth inverted over mercury. If it be exposed to the air, after this process, for ever so short a time, it immediately absorbs some air; a proof of the strong affinity which it has for that fluid *. This philosopher has pointed out the following method of ascertaining whether water be perfectly free from air. Tinge the water blue with litmus, fill a flask with it, invert the flask under water, and introduce into it pure nitrous gas till about to of the vessel is filled. If the water contain air, a portion of the nitrous gas will combine with its oxygen, and be converted into nitric acid. The consequence of which will be, that the litmus will assume a red colour. Even the proportion of air in the water may be estimated by the quantity of ammonia necessary to restore the blue colour to the litmus. If the litmus retains its blue colour, we may be certain that the water which we try contains no sensible portion of air +. It is proper to remark, howeyer, that this experiment does not succeed in a satis-

WATER.

nanner, unless an unusual quantity of oxygen present. No effect is produced when nitrous itated with good spring water.

ater has no action on the simple combustibles Action of ld, nor does it combine with any of them. combustiin does not act upon it even at a red heat; but at that temperature decomposes it, forming acid, and various species of heavy inflamma-

the simple

The action of phosphorus at a red heat has Sulphur, as far as is known at present, decompose it.

the metals, iron, zinc, antimony, and tin, de- Of the meit when assisted by heat; silver, gold, copplatinum, have no effect upon it. The action ther metals at a red heat has not been tried.

ater dissolves the alkalies and alkaline earths. ths proper are insoluble in it. It dissolves also I salts, and is capable of combining with a great of bodies.

and earths.

ater has the property of uniting with bodies in Water comrent ways. Some it dissolves; the compound forms. liquid, like water. In this way it unites with salt, with sugar, and a vast number of other Other substances combine with it without loir solidity. The water in this case loses its lim, and assumes that of the substance to which In this way it unites to lime, to alumiany saline bodies, and to a variety of metallic

When the compound of water with another e remains liquid, the proportion of water is un-

[#] Mem. Par. 1781. p. 273.

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Book II. Division II. limited; but when the combination formed is solid, the water combines always in a certain determinate proportion.

To the first of these combinations, or the solutions of bodies in water as they have been termed, considerable attention has always been paid; but the second kind, in which the water assumes a solid form, was very much overlooked, if not altogether neglected, till the attention of chemists was drawn to them by Mr Proust, who has given to such combinations the name of bydratu. Thus the combination of lime and water, usually called slacked lime, is in his language a bydrate of lime: in like manner, the crystals of barytes and strontian are hydrates of these alkaline earths, and crystallized potash and soda are hydrates of the fixed alkalies. Though the term bydrate is in some respects exceptionable, we shall continue to use it, as no other has been suggested. The compounds called bydrates deserve attention, since they differ very considerably in their properties from those bodies with which they have been hitherto confounded. Thus the hydrates of lime, barytes, potash, &c. are very different in several respects from lime, barytes, potash, &c.

Alkaline and earthy hydrates The hydrates of potash and soda are crystallized, and contain always a determinate proportion of water: according to Proust, about 30 per cent. When dissolved in water they occasion coid; whereas the alkalies is the state of powder produce heat. The same observations are applicable to the hydrates of barytes and strontian. The proportion of water which combines with lime is much less considerable than that which enters into the composition of the hydrates just mentioned. Hence the hydrate of lime does not crystallize, but re-

Hydrates.

Chap. I.

mains usually in the state of a powder. We know, however, that it retains the water with great obstinacy, and that the formation of it is one of the principle causes of the solidification of mortar *. The hydrate of alumina is no less remarkable for the obstinacy with which it retains its water. It is the substance to which Saussure gave the name of spongy alumina.

But the hydrates of the metallic oxides are the most Metalline. remarkable. They first drew the attention of Proust, and induced him to invent the term hydrate. His observations, as far as regards the bydrate of copper, have been called in question by Berthollet junior +; but his arguments do not appear to me sufficiently conclusive to overturn the ingenious theory of the Spanish chemist. He has demonstrated indeed, that the bydrate of copper usually retains a small portion of acid; but he has not shown that its peculiar properties are owing to that acid, while Proust has made the contrary more than probable, by ascertaining that the properties of the hydrate remain the same, even when the proportion of acid varies, and when it is so far diminished as to be no longer perceptible.

When copper is dissolved in nitric acid, and a suffi- Hydrate of cient quantity of potash added to the solution, a blue powder falls to the bottom, which when sufficiently washed, and carefully dried, coheres together, and forms a brittle mass breaking with a vitreous fracture. This mass is the hydrate of copper. It has an exquisitely disagreeable taste, and acts with great energy upon the

copper.

[#] See Proust, Jour. de Phys. lix. 347.

[†] Statique Chemique, ii. 455.

Pook II.
Division II.

system when swallowed, or even kept in the mouth. When distilled it yields 25 parts of water, and leaves 75 of black oxide of copper.

Of iron,

When the solution of iron in sulphuric acid is treated in the same way, a green powder falls, which is a hydrate of iron. These two hydrates are delicate, and easily lose their water. But some of the other hydrates of the metallic oxides retain their water with great obstinacy. This is the case particularly with the hydrates of nickel and cobalt, which resist the action of a considerable heat.

Of tip.

When tin dissolved in muriatic acid is precipitated by potash, a white powder is obtained, which washed and dried in the heat of boiling water is a hydrate of tin. When distilled in a retort it loses 5 per cent. of water, and is converted into protoxide of tin †.

Most of the metallic hydrates are remarkable for the brilliancy of their colours. They are much more easily dissolved by acids than the oxides; and when put into the mouth they affect the organs of taste even more powerfully than the metallic salts.

Gases con-

- 10. All gases in their usual state contain combined with them a quantity of water, which often amounts to a considerable proportion of their weight. Part of this water may be abstracted by exposing the gases to substances which have a strong affinity for water, as dry potash; but part adheres with a great deal of obstinacy, and perhaps cannot be removed by any method in our power.
 - 11. Water was believed by the ancients to be one of

Proust, Jour. de Bbys. lix. 347.

the four elements of which every other body is composed; and, according to Hippocrates, it was the substance which nourishes and supports plants and animals. That Opinions awater was an unchangeable element continued to be be- ture of walieved till the time of Van Helmont, who made plants grow for a long time in pure water: from which experiment it was concluded, that water was convertible isto all the substances found in vegetables. Mr Boyle having digested pure water in a glass vessel hermetically sealed for above a year, obtained a quantity of earthy scales; and concluded, in consequence, that he had converted it partly into earth *. He obtained the same earth by distilling water in a tall glass vessel over a slow fire +. Margraff repeated the expetiment with the same result, and accordingly drew the same conclusion. But the opinion of these philosophers was never very generally received 1. The last person who embraced it was probably Mr Waselton, who published his experiments on the subject in the lournal de Physique tor 1760. Mr Lavoisier had proved, as early as 1773, that the glass vessels in which the distillation was performed lost a weight exactly count to the earth obtained. Hence it follows irresistibly that the appearance of the earth, which was silica, proceeded from the decomposition of the vessels; for glass contains a large proportion of silica. It has been since shown by Dr Priestley, that water always decom-

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⁺ Ibid. i. \$67. Shaw's Boyle, ill. 417. t we an account of all the facts relating to this subject in Watson's Chemical Estage 18- 257.

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Division II.
History of the discovery of its component parts.

poses glass when applied to its surface for a long time in a high temperature.

Water is now known to be an oxide of hydrogen, or a compound of oxygen and hydrogen. As this discovery has almost entirely altered the appearance of the science of chemistry, by furnishing an explanation of a vast number of phenomena which were formerly inexplicable, it will be worth while to give a particular account of the different steps which gradually led to it.

The first person probably who attempted to discover what was produced by burning hydrogen gas was Scheele. He concluded, that during the combustion oxygen and hydrogen combined, and that the product was caloric.

In 1776 Macquer, assisted by Sigaud de la Food, set fire to a bottle full of hydrogen gas, and placed a saucer above the flame, in order to see whether any faliginous smoke would be produced. The saucer remained perfectly clean; but it was moistened with drops of a clear liquid, which they found to be pure water.

Next year Bucquet and Lavoisier exploded oxygen and hydrogen gas, and made an attempt to discover what was the product; about the nature of which they had formed different conjectures. Bucquet had supposed that it would be carbonic acid gas; Lavoisier, on the contrary, suspected that it would be sulphurie or sulphurous acid. What the product was they did not discover; but they proved that no carbonic acid gas was

Macquet's Dictionary, art. Gas inflammable.

ed, and consequently that Mr Bucquet's hypotheras ill founded †.

Chap. I.

the beginning of the year 1781, Mr Warltire, at equest of Dr Priestley, fired a mixture of these two s contained in a copper vessel; and observed, that the experiment the weight of the whole was dished. Dr Priestley had previously, in the presence Ir Warltire, performed the same experiment in a vessel. This vessel became moist in the inside, was covered with a sooty substance ‡, which Dr stley afterwards supposed to be a part of the merused in filling the vessel §.

the summer of 1781, Mr Cavendish, who had informed of the experiments of Priestley and Itire, set fire to 500,000 grain measures of hydrogas, mixed with about 2½ times that quantity of mon air. By this process he obtained 135 grains of water. He also exploded 19,500 grain measures tygen gas, with 37,000 of hydrogen gas, and obed 30 grains of water, containing in it a little nitric. From these experiments he concluded that water

compound.—Mr Cavendish must therefore be coned as the real discoverer of the composition of wa-He was the first who ascertained that water is uced by firing oxygen and hydrogen gas, and the

who drew the proper conclusion from that fact. Watt, indeed, had also drawn the proper conclufrom the experiments of Dr Priestley and Mr : ltire, and had even performed a number of experi-

[†] Mrm. Par. 1981, p. 470.

f Priestley, v. 395.

³ Phil. Trans. laxiv. 332.

Book II. Division II. ments himself to ascertain the fact before Mr Caverdish had communicated his; but he had been deterred from publishing his theory by some experiments of Dr Priestley, which appeared contrary to it. He has therefore a claim to the merit of the discovery; a claim however, which does not affect Mr Cavendish, who knew nothing of the theory and experiments of that ingenious philosopher.

Meanwhile, in the winter 1781-2, Mr Lavoisier, who had suspected that when oxygen and hydrogen gas are exploded, sulphuric or sulphurous acid is produced_ made an experiment in order to ascertain the fact, at which Mr Gengembre assisted. They filled a bottle. capable of holding six pints (French), with hydrogen gas, to which they set fire, and then corked the bottle, after pouring into it two ounces (French) of lime-water. Through the cork there passed a copper tube, by means of which a stream of oxygen gas was introduced to support the flame. Though this experiment was repeated three times, and instead of lime water a weak solu tion of alkali and pure water were substituted, the could not observe any product whatever +. sult astonished Mr Lavoisier exceedingly: he resolved = therefore, to repeat the experiment on a larger scale and if possible with more accuracy. By means of pipe furnished with stop-cocks, he put it in his power terms supply both gases as they should be wanted, that he might be enabled to continue the burning as long as here thought proper.

The experiment was made by Lavoisier and La Plac =

[◆] Bil. lxxv. 330.

the 24th of June 1783, in the presence of Messrs Roi, Vandermonde, several other academicians, and Charles Blagden, who informed them that Mr Cawish had already performed it, and that he had obmed water . They continued the inflammation till their stock of gases was wasted, and obtained about grains of water, which, after the most rigid examition, appeared to be perfectly pure. From this exment Lavoisier concluded, that water is composed baygen and hydrogen. Mr Mongez soon after permed the same experiment, and obtained a similar re-16: and it was repeated again by Lavoisier and Meuson a scale sufficiently large to put the fact beyond abt †.

drogen is, that when these two gases, mixed in proproportions, are fired, they almost wholly disapand there is found in their place a quantity of water, as nearly equal to them in weight as can expected in experiments of that delicate nature. The frogen gas is made to pass slowly from the glass jar which it is contained, by means of a tube furnished a stop-cock, into a glass globe filled with oxygen It is set on fire at the extremity of the tube, eiby means of electricity or by a little phosphorus. It continues to burn slowly till the whole of it is nmed. New portions of oxygen gas are introduoccasionally from another glass jar, by means of a

furnished with a stop-cock. The water, as it is

med, is condensed in the glass globe. A great num-

The proof that water is a compound of oxygen and Proof.

t Ibid p. 474

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· Mem. Par. 1781, p 472

Book II. Division II. ber of precautions are necessary to ensure the purity of the gases, and to measure their weight and the nature of the gas which remains after combustion. But for these I refer to the account of the experiments themselves, which have been published by the French chemists in the Memoirs of the Academy of Sciences. The experiment on which the greatest dependence may be put was made in the year 1790 by Seguin, Fourcroy, and Vauquelin*. The bulk of the gases employed in this experiment was,

French Inches.

Hydrogen gas.....25980.563

Oxygen gas.....12479.080

Total.....38459.643

The water obtained amounted to 7249.227 grains rench, or 3947.3 grains troy, or 12.390 oz. It exhibited no mark of acidity, and appeared in every respect to be pure water. Its specific gravity was to that of distilled water as 18071 to 18630; or nearly as 1.000053 to 1.

The residuum of gas in the vessel after combustion amounted to 987 cubic inches French; and, on being examined, was found to consist of the following quantities of gases:

Sec Ann. de Chim. viji. 230

WATER.

	French Inches.
Azatic gas	467
Carbonic acid gas	39
Oxygen gas	465
Hydrogen gas	16
Total	987
Now the weight of the v	vhole
gases employed was	6306.71
That of the water obtained,	, and
of the residuum	6303.24
	مسينيه مسييه
^	_

grains less than had been employed. This approaches as near an equality as can be expected in experiments of this nature. The small surplus of azotic gas found after the combustion cannot be accounted for, unless we suppose some common air to have gained admission during the process.

As sufficient precautions had been taken to prevent the introduction of earbonic acid gas, the quantity found in the residuum must have been formed during the process. There must therefore have been a small quantity of carbon introduced. Now zinc, it is supposed, often contains carbon, and hydrogen has the property of dissolving carbon; probably, then, the carbon was introduced in this manner. The carbonic acid found in the residuum amounted to 23.306 grains, which according to Lavoisier's calculation, is composed of 8.256 grains of carbon and 14.348 grains of oxygen:

Subtracting these 8,958 grains of curbon, and the

Book II. Division II. 0.530 of a grain of hydrogen which remained in the vessel, from the total of hydrogen introduced, there will remain 852.600 grains for the hydrogen that disappeared.

Subtracting the 14.348 grains of oxygen which entered into the composition of the carbonic acid, and the residuum of oxygen, which amounted to 188.371 grain, the quantity of oxygen that disappeared will amount to 5094.6 grains.

Which is less than the agrains gases consumed by

It is impossible to account for the exact coincidence of the water condensed with the weight of the gases consumed, unless we suppose it to be composed of the bodies.

This experiment gives us the composition of water
as follows:.....Oxygen...85.662 grains troy
Hydrogen 14.338

100.000

Dr Priestley, however, who made a great many experiments on this subject, drew from them a very different conclusion; and thought he had proved, that during the combustion the two gases combine, and that the combination is nitric acid. This theory was adopt-

or rather it was suggested, by Mr Keir, who has sported it with a great deal of ingenuity ".

Chap, I.

Let us examine these experiments of Dr Priestley +. dee whether they warrant the conclusions he has wn from them. The gases were exploded in vessels copper. He found that the quantity of water obed was alway less than that of the gases which he a used. He obtained also a considerable quantity of In the experiment made on the largest ric acid. antity of the gases, and from which he draws his eclusions, the quantity of liquid obtained amounted 42 grains. This liquid was examined by Mr Keir. was of a green colour; 72 grains of brown oxide of per were deposited in it, and it contained a solution mitrate of copper (copper combined with nitric acid). Keir analysed this liquor: It consisted of pure waand nitrate of copper: and Mr Keir concluded that mitric acid formed amounted to stath of the oxygen employed. Here then a quantity of oxygen and drogen gas has disappeared: What has become of em? They have combined, says Dr Priestley, and med nitric acid. This nitric acid is only to their might. Dr Priestley supposes, however, that it conhas the whole oxygen and hydrogen that existed in gases, and that all the rest of the weight of these was owing to a quantity of water which they had in solution. Oxygen gas, then (for we shall neet the hydrogen, which Dr Priestley was not able to into view at all), is composed of one part of oxyand 19 of water. Where is the proof of this? Dr they informs us, that he ascertained by experiment

Kent's Distinuery, art Nurvis Acid. 4 Phil. Trans. 2788.

that half the weight of carbonic acid gas was pure we ter. Supposing the experiment accurate, surely it can not be concluded from it that oxygen gas consists c To parts, or almost wholly of water. It is impossible therefore, from Dr Priestley's experiments, allowing hi ingenious suppositions and conjectures their atmos force, to account for the disappearing of the two gasts, or the appearance of the water, without admitting that this liquid is actually composed of oxygen and hydrogen. If we add to this, that oxygen gas can scarcely be procured absolutely free from some mixture azote, and that his oxygen was always obtained eithe from red oxide of lead, or from black oxide of mange nese, or red oxide of mercury, all of which substances yield a considerable proportion of azote; if we add, the it has been proved beyond the possibility of doubt, and to Dr Priestley's own satisfaction, that nitric acid composed of oxygen and azote—we shall find it no di= ficult matter to explain the origin of that soid in IX Priestley's experiments: and if we recollect that in Se gnin's experiment, upon a much larger scale than D Priestley's, no nitric acid at all was formed, it will be impossible for us to believe that the compound forme. by exygen and hydrogen is nitric acid. Priestley's experiments rather confirm than destroy the theory of the composition of water. We obtain from them, however, one curious piece of information, that the presence of copper increases the quantity of nitriacid formed.

The proof for the composition of water, derived from the combustion of hydrogen gas, is rendered still strong er by reversing the experiment. When electric explosions are made to pass through water, part of it is decomposed and converted into anygen gas and hydrogen

Chap. L

Messrs Van Troostwyck and Dieman, assisted by inthertson, filled a small class tube, ! th of an inch meter a 112 inches long, with distilled water. and of this tube was sealed bermetically, but at me time a small gold wire had been passed through Snother wire passed through the open end of the and could be fixed at greater or smaller distances the first wire. By means of these wires, they great number of electrical explosions pass through Bubbles of air appeared at every exploind collected at the top of the tube. When elecbarks were passed through this air, it exploded lappeared almost completely. It must therefore consisted of a mixture of oxygen and hydrogen and this gas must have been formed by the decomon of the water; for they had taken care to dethe water beforehand of all its air, and they used precaution to prevent the access of atmospherical and, besides, the quantity of gas produced did not ash, but rather increase, by continuing to operate ber of times upon the same water, which could we been the case had it been merely air dissolved ter: nor would atmospherical air have exploded, hit only a very small residuum, not more than yoth

They had taken care also to prove that the elecpark did not contribute to form hydrogen gas; for asing it through sulphuric and nitric acids, the let was not hydrogen, but oxygen gas.

on, assisted by Mr Cuthbertson. He produced, cans of electricity, quantities of gas from water, nating to 56.5488 cubes of 1's th of an inch each; on

[.] Jour. de Phys. xxxv. 369.

nitrous gas being added to which, it suffered a diminution of bulk, and nitrous acid appeared to have been formed. It must therefore have contained oxygen gas. When oxygen gas was added to the remainder, and an electric spark passed through it, a diminution took place precisely as when oxygen and hydrogen gas are mixed: It must therefore have contained hydrogen. When an electric spark was passed through the gas thus produced from water, the gas disappeared, being, no doubt, converted into water *.

Such are the proofs by which the component parts of water have been ascertained. If we consider them attentively, and compare them with a vast number of other chemical phenomena, all of which tend to confirm and establish them, we must allow, I think, that scarcely any physical fact whatever can be produced, which is supported by more complete evidence. There are indeed some galvanic phenomena which scarcely seem compatible with it; but the nature of this singular power is still too imperfectly understood to warrant even a conjecture concerning it.

II. COMBUSTIBLE OXIDES.

THE oxides formed by all the simple combustibles, except hydrogen, are combustible, and of course cannot be formed by combustion. The composition of these oxides is still imperfectly known, owing to the extreme difficulty of examining them.

^{*} Nicholson's Jour. i. 242. These experiments are now made with great case by the galvanic apparatus.

Chap. I.

SECT. II.

OXIDES OF CARBON.

It is an opinion at present pretty generally admitted by chemists, that carbon is capable of uniting with at least two doses of oxygen, and of forming two compounds, one of which is an oxide, and one an acid. The oxide is carbonic oxide, which is a combustible gas. A short ketch of the properties of this oxide has been given in preceding part of this Work. But it will be necesmary here to enter more into detail. Besides carbonic oxide, it was supposed, from the experiments of Morweau on the diamond, that there existed another oxide of carbon, containing a smaller proportion of oxygen, and that this oxide was nothing else than pure charcoal. But the late experiments of Messrs Allen and Pepys, by demonstrating the inaccuracy of Morveau's experiments, have destroyed the evidence upon which that supposition was founded. We are at present ignorant of the composition of charcoal, though it has been demonstrated that it contains at least two ingredients, carbon and hydrogen. Till the composition of this substance be better ascertained, I shall allow it to retain the place in the arrangement of chemical substances which was assigned it when it was considered as an oxide of carbon.

I. OF CHARCOAL AND CARBONOUS OXIDE.

BEFORE the experiments of Morveau on the diamond were made known, chemists were accustomed to confound together carbon and ebarcoal, though they had been carefully distinguished by Lavoisier who indeed invented the term carbon, to render it more difficult to mistake for charcoal, the substance to which he applied it. After these experiments became known, the mists fell into the new mistake of confounding charcoal with carbonous oxide, till this error was in some measure rectified by Cruickshanks and Berthollet, and till the experiments of Allen and Pepys demonstrated in inaccuracy.

Two species of charcoal, commen 2. When charcoal is prepared in the usual way, by exposing wood in close vessels to a red heat, it shways contains a portion of hydrogen: For if a quantity of this charcoal be exposed to a strong heat in a retort of posselain, iron, or coated glass, a great quantity of gur is obtained. The gas which comes over first is a mixture of carbonic acid and heavy inflammable gas; but the proportion of carbonic acid diminishes, and at last it ceases to come over at all; yet the inflammable gas continues as copious as ever.

The evolution of these gases was long ascribed by chemists to the water which charcoal usually contains, and which it is known to absorb from the atmosphere with considerable avidity. If that were the case, the proportion of inflammable gas ought to diminish at the same rate with the carbonic acid; the hydrogen of the

^{*} Cruickshauks, Nicholson's Jenrnal, 1802. v. 210.

one being equally derived from the decomposition of Chap. L. water with the oxygen of the other. But as the evolution of inflammable gas continues after that of carbonic acid has ceased, it is scarcely possible to deny, that the hydrogen which thus escapes constituted a component part of the charcoal.

2. When common charcoal is exposed for an hour And prepain a close crucible to the strongest heat of a forge, it ceases to emit gas, and no temperature is sufficient to expel gas from charcoal thus treated . Desormes and Clement have endeavoured to demonstrate, that by this breatment common charcoal is deprived of the whole of its hydrogen. They put a quantity of charcoal, recently exposed to the heat of a forge, and not yet cold, Into the middle part of a long glass tube. To each extremity of the tube was fixed another tube, filled wift dry muriate of lime, and surrounded with a mixture of snow and salt. To the extremity of one of these tubes was fixed an empty bladder; to the extremity of the other, a bladder containing a quantity of oxygen gas. The charcoal was heated to redness by placing the tube in a furnace, and then the oxygen gas was made to pass slowly over it from one bladder to the other. By this contrivance the charcoal was burnt, and converted into carbonic acid. The oxygen gas, in passing through the muriate of lime, was made to deposite the meisture which it contained, and the proportion was known by the increase of weight of the muriate. If the charcoal contained hydrogen, water would be formed during its combustion, which would unite with the

Desormes and Clement, Aun. de Chim. xxxix. 29.

carbonic acid formed. But this gas would deposite its moisture during its passage through the muriate of lime, at the other extremity of the tube, and the increase of weight which this muriate would experience would indicate the proportion of water formed during the process. In both cases the muriate of lime was increased in weight 0.02 parts. Were we to suppose this increase owing to the formation of water, the small quantity would only contain hydrogen to the amount of Tree of the charcoal; a portion too small to be regarded *.

The same chemists tried the combustion of charcoal obtained from a variety of other substances exposed to the heat of a forge, as pitcoal, animal substances, and various vegetable substances, and found the products exactly the same. Hence they conclude that charcoal is in all cases the same, provided it be exposed to strong enough heat. And they conclude, too, that by this strong heat the whole hydrogen of common charcoal is expelled.

But this is going rather farther than their experiments will warrant; and it is directly contrary to the experiments of Cruickshanks, who always found gases obtained by means of charcoal, in whatever state, to contain hydrogen. Besides, Berthollet has pointed out circumstances which render the precision of these chemists somewhat doubtful.

Both contain some hydrogen. 3. Such are the facts at present known respecting the composition of charcoal. They enable us to conclude, that there are two species of charcoal, namely, common

Ann. de Chim. xlii. 128.

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and prepared charcoal. The first contains at least two ingredients, carbon and hydrogen; the second is deprived of a portion of its hydrogen. It consists chiefly of carbon; but it still retains a small portion of hydrogen, and is not, therefore, strictly speaking, pure carbon.

4. When the diamond is exposed to the action of heat and air, it has been observed by different experimenters to acquire a black coat not unlike charcoal. This coat may be considered as a combination of carbon and oxygen; it is perhaps carbonous oxide in a state of purity. Whether any similar combination exists nasive has not yet been ascertained.

II. OF CARBONIC OXIDE.

The substance at present known by the name of car- History. donic oxide, is a gas which had been confounded with carbureted hydrogen, till Dr Priestley drew the attention of chemists to it in a dissertation which he pubished in defence of the doctrine of phlogiston. His experiments were immediately repeated, and his opiaions confirmed by Dr Woodhouse of Pennsylvania. But the real nature and composition of the gas was discovered by Mr Cruickshanks of Woolwich, and the discovery communicated to the public in 1802". About the time of the publication of Mr Cruickshanks' dissertation, the experiment of Woodhouse, which he had transmitted to France, engaged the attention of the National Institute. Guyton Morveau, who had been

Nicholson & Journal, v. 1. and 201.

appointed to give a detailed account of these experiments, engaged Clement and Desormes to investigate the subject; and these gentlemen were gradually led to the same conclusions * which had been previously drawn by Cruickshanks. The subject in the mean time attracted the attention of Bertholet, whose experiments induced him to form a different opinion respecting the composition of charcoal and carbonic oxide from that which had been entertained by the other chemists. This opinion he supported in three elaborate dissertations, published n the fourth volume of the Memoirs of the National Institute; in which he examines the experiments, and combats the conclusions of the other chemists, with his usual sagacity. About the same time a dissertation was published by the Dutch chemists, contradicting the experiments of all other philosophers, and affirming the real results to be very different †. This short historical sketch is a sufficient proof of the great difficulty attendng the investigation. No less than four different opinions have been maintained, and every one of them by men of eminence, of acknowledged skill, and undoubted candour.

Prepara-

1. There are four different processes by which carbenic oxide gas may be procured: First, When a mixture of purified charcoal and the oxides of iron or zinc, or indeed of any oxide capable of bearing a red heat, is expessed to a strong heat in an iron retort, the oxide is gradually reduced, and during the reduction a great quantity of gas is evolved. This gas is a mixture of carbonic acid gas and another which burns with a blue flame. It is to

Ann. de Chim. Exxix. 88. and zlii. 121.

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last that the term carbonic oxide has been applied. the carbonic acid may be separated by passing the gas rough lime-water. Mr Crnikshanks tried in this way e oxides of iron, zinc, and copper, litharge, and the lack oxide of manganese. The following conclusions bult from his experiments: Those oxides which part with their oxygen most readily yield the greatest proprtion of carbonic acid; those that retain their oxyen most obstinately yield the greatest proportion carbonic oxide. It is always towards the begining of the process that the greatest proportion of carmic acid gas comes over; it gradually diminishes, at last nothing but carbonic oxide is disengaged *. The results obtained by Clement and Desormes coinde almost exactly with the experiments of Cruikshanks. Hot they satisfied themselves with the white oxide of inc, without trying those of other metals. They sublitured plumbago for charcoal, and obtained the same tsuits +.

Second, When a mixture of one part of purified charal, and three parts of the carbonate of lime, or of stronam, or of barytes, is exposed to a strong heat in an iron wort, the carbonic acid is gradually separated or deempased, and gas is evolved to abundance. This gas maists of a mixture of about one part of carbonic acid five parts of carbonic oxide 1. In this case a poron of the carbonic acid of the carbonate is disengaged caltered, but the greatest part of it is converted into arbonic oxide by the action of the charcoal.

Nicholson's Jour. 1801, v. 2.

Ann. de. Chim. xxxix. 33.

Third, When a mixture of equal parts of any of the three above-mentioned earthy carbonates and clean iron filings is heated strongly in an iron retort, the carbonic acid is decomposed in like manner by the action of the iron, and the very same gases are procured in great abundance. Dr Priestley first tried this method with the black oxide of iron and carbonate of barytte; but when Cruickshanks substituted pure iron, the gaseous product was considerably increased.

Fourth, When carbonic acid gas is made to pass slowly and repeatedly through prepared charcoal heated to redness in a porcelain or iron tube, it gradually disappears, and carbonic oxide is found in place of it. Here the charcoal decomposes the carbonic acid precisely as in the two last cases, with this difference only, that it is in a gaseous state, whereas in them it was combined with a base. This experiment was first made by Cruickshanks †, and afterwards by Clement and Desormes ‡.

How purified.

2. Such are the different processes for procuring carbonic oxide. From the experiments of Cruikshanks we learn, that the third method is the only one to be depended on for obtaining the gas in a state of purity. If equal parts of chalk and iron filings, previously exposed to a red heat separately in close vessels, be mixed together and strongly heated in an iron retort, the gases which come over are merely a mixture of carbonic acid and carbonic oxide; and the first being abstracted by

^{*} Nicholson's Jour. 1802. v. 4. and 208. † Ibid. p. 209.

Ann. de Cb m. xxxix. 46.

of lime-water, the carbonic oxide gas remains in b of purity.

Carbonic oxide gas, thus obtained, is invisible and Properties. is like common air. Its specific gravity, according experiments of Cruickshanks, is 0.956, that of ing 1 000. It is to common air as 22 to 23. One red cubic inches of it weigh 30 grains. The repotained by Desormes and Clement does not differ from this ".

simals cannot breathe this gas without suffocation. put into it by Desormes and Clement dropt down before they had time to take them out; and when attempted to breathe it themselves, the consequence kiddiness and faintness +. Neither will any comthe body burn in it.

is not altered by exposure to light, nor by passing cough a red hot tube. From the experiments of ent and Desormes, we find that it is dilated by exactly like common air, as was indeed to be ex-

Carbonic oxide gas is combustible. It takes fire Combust:open air when it comes in contact with a red hot ble. or when it is presented to the flame of a candle, forms with a lambent blue flame. When mixed common air before it is kindled, it burns more raand brilliantly, but does not detonate. The com-

taking a mean of their experiments, we obtain the weight of a 1 1 35 grammes, which reduced to our standard gives the weight cubic inches 28 7 grains. Their result would have been still Eruikshanks', had I excluded their trials with some of the lightimpurest gases which they obtained.

de Char. 22212.56.

bustion is still more rapid and brilliant if we substitute oxygen gas for common air. Sometimes it detonates with oxygen, but most commonly the detonation does not take place.

From the experiments of Cruickshanks, which have been confirmed by those of Clement and Desormes, we learn that 160 cubic inches of carbonic oxide, in order to undergo complete decomposition, must be mixed before combustion with 40 cubic inches of oxygen gas; and this mixture, being exploded by electricity in adetonating tube, is converted into 92 cubic inches of carbonic acid gas. Or, in numbers, 30 grains of carbonic oxide require for saturation about 13.6 grains of oxygen; and the resulting compound amounts to 43.6 grains of carbonic acid gas. No sensible quantity of water is formed by the combustion of this gas when it is procured by the third process, and made as dry and pure as possible before the experiment.

Action of simple combustibles, 5. This gas has no action whatever upon the simple combustibles at the common temperature of the atmosphere; but its activity is somewhat augmented by the assistance of heat.

When passed through melted sulphur, it does not combine with it, nor alter its properties; but it dissolves a little phosphorus, and acquires the property of burning with a yellow flame. When passed through red hot charcoal, it dissolves a part of it, if we believe Desormes and Clement, and its specific gravity is increased. The same chemists affirmed, that when a mixture of carbonic oxide and hydrogen gas is made to pass through a red hot glass tube, charcoal is deposited, which lines the inside of the tube with a shining enamel; that water is formed, and hydrogen, seemingly pure.

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disengaged from the other end of the tube *. But when this experiment was repeated by Saussure junior, he found that the supposed enamel of charcoal was merely the black (or rather bluish) colour which flint glass acquires when hydrogen is brought in contact with it at a red heat, as had been previously observed by Dr Priestley †. Indeed it is very unlikely that hydrogen gas is capable of decomposing carbonic oxide; as Saussure has shown, that when carbonic acid and hydrogen gas are made to pass through a red hot tube, the acid is decomposed, and carbonic oxide produced—a result which has been even confirmed by the subsequent experiments of Clement and Desormes.

6. None of the simple incombustibles produce Of incomany change upon carbonic oxide at any temperature hitherto tried; but the action of oxymuriatic acid gas upon it is extremely curious and important. investigation of this part of the subject we are entirely indebted to Mr Cruikshanks, who has thereby disclosed to chemists a new and valuable method of trying the purity and composition of the combustible gases:

If a phial be filled with a mixture of two measures of carbonic oxide gas and $2\frac{3}{1}$ measures of oxymuriatic acid gas I, then closed with a ground stopper, and allowed to remain for 24 hours with its mouth inverted under mercury, on drawing the stopper under water, two-thirds of the gas are immediately absorbed, and all the rest by agitation in lime-water (except th of a

Of oxymuriatic seid.

Ann. de Chim. xxxix. 61. † Jour. de Phys. lv. 396.

[#] Procured by pouring muriatic acid on the hyperoxymuriate of pot-**23h.**

measure of azote). Hence we see that these two gases act upon each other at the temperature of the atmosphere; that the carbonic oxide gradually abstracts oxygen from the other gas, and is converted into carbonic acid. The oxymuriatic gas thus decomposed is converted into muriatic acid, which is instantly absorbed on the admission of water. Thus by the mutual action of the two gases, the whole is converted into carbonic acid and muriatic acid.

Mr Cruikshanks ascertained that this mixture of carbonic oxide and oxymuriatic acid gas does not bum when electric sparks are made to pass through it, nor is its nature altered; whereas a mixture of carbureted hydrogen and oxymuriatic acid gas explodes immediately. This difference enables us to distinguish carbureted hydrogen from carbonic oxide with the greatest facility †.

Of metals,

7. From any experiments hitherto made, it does not appear that carbonic oxide is capable of acting on the metals; but Clement and Desormes affirm, that when passed hot over the red oxide of mercury, it produces a commencement of reduction. Indeed it is very likely that it will be found capable of reducing several of the metallic oxides, especially those that part with their oxygen easily.

Alkalies
and earths.

8. Neither the fixed alkalies nor the earths have any action on carbonic oxide. Neither does ammonia alter it when passed with it in the state of gas through a red hot tube ‡.

^{*} Nicholson's Journal, 1802, v. p. 205. † Ibid. p. 207.

[†] Clement and Desermes, Ann. de Chim. xxxix. 62.

9. From the history of the properties of this gas just given, it must be obvious at once, that it contains car- Composibon as an ingredient; for when fired with oxygen, it yields carbonic acid gas as a product. That it differs from carbureted hydrogen is obvious from its specific gravity, from the action of oxymuriatic acid gas, and from the result obtained by burning it. As it yields on combustion no perceptible portion of water, it was concluded by Cruikshanks, and afterwards by Guyton Morveau, Desormes and Clement, that it contained no hydrogen, and contained no other combustible base besides carbon. But it requires much less oxygen gas for combustion than charcoal. Thus 100 parts of charcoal require 257 parts of oxygen to saturate them, whereas 100 parts of carbonic oxide require only 454 parts of oxygen; and in both cases carbonic acid is produced. This remarkable difference can only be accounted for by supposing that the carbonic oxide is already combined with a portion of oxygen, and therefore a smaller addition must be sufficient to saturate it. Accordingly this was the consequence drawn by Cruikshanks; and it is impossible to resist the evidence in favour of his conclusions. The gas therefore, according to this reasoning, is a compound of carbon and oxygen. Hence the name carbonic oxide given to it by chemists.

If we suppose, according to the experiments of Lavoisier, that carbonic acid is composed of 28 carbon and 72 oxygen, and consider the experiments of Cruikshanks as approaching to accuracy, it will be easy for us to ascertain the component parts of this oxide.

According to Cruikshanks, 30 grains of carbonic oxide combine by combustion with about 13.6 grains of

7: /

bent 43.6 grains. Hence it follows, that carbonic acid is composed of about

69 carbonic oxide 31 oxygen

100

But 100 carbonic acid are composed of 72 oxygen and 28 carbon. We have therefore this equation,

Carbon. Oxygen. Carb.Ox. Oxygen.

28 + 72 = 69 + 91

Carbon. Oxygen. Carb. Oxide.

26 + 41 = 69

That is to say, 19 parts of carbonic oxide are composed of 28 parts of carbon and 41 of oxygen. Of course we have fer cent. about

41 carbon

59 cxygen

100 carbonic oxide

And 100 ports of carbon united to 146 of oxygen form

Throwald. Himbolies. But all this statement has been called in question by Bert'o'. According to this sagacious philosopher, there are two different species of inflammable gases containing carbon. The first species is composed of eathon and hydrogen; the second, of carbon, hydrogen, and oxygen. To distinguish these two species from tach other, he calls the first carriar and hydrogen; the second extraordal hydrogen. To the first species belong the gases obtained by passing alcohol through a red hot tube, by distilling oil, and by exposing moist charcoal to a red h.a., &c. To the second species belong the gases where it is a first species belong the gases where it is a first species belong the said has a red hot sube, by distilling oil, and by exposing moist charcoal to a red h.a., &c. To the second species belong the said housel by exposing contract to a strong

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heat, the gas obtained by distilling sugar, the gas described in this occor under the name of carbonic oxide, &c. There are many varieties of these gases, differing from each other in the proportions of their ingredients: and the gases belonging to the first species may, by various processes, be converted into the second. He calculates the proportion of hydrogen in carbonic oxide to be about the of the whole.

These conclusions are obviously irreconcileable with the experiments of Cruikshanks and of Desormes and Clement, and cannot therefore be admitted without very decisive proofs. The carbonic oxide obtained by Cruikshanks, when as pure as possible, yielded no perceptible quantity of water when burnt with oxygen. But Berthollet observes that all gases contain water as a constituent part, and shows that all the water that could have been formed would have combined with the carbonic acid gas, and remained invisible. Even if we were to grant this, and to allow that some water has been formed in every case of the combustion of carbonic oxide with oxygen hitherto tried, still Mr Berthollet's hypothesis would stand upon as weak ground as ever. For it is obvious from the experiments of Cruikshanks, that the drier he made his materials the purer was the gas, or the less water was formed during its combustion; and when he employed bodies that could not contain any hydrogen except in , the state of water (namely, dry chalk and tin or iron), the gas was obtained purest of all.

Mr Berthollet's chief reason for affirming that carbonic oxide must contain hydrogen is its small specific gravity. The specific gravity of

oxygen gas is 1.103

OXIDES

Book II. Division II. carbonic oxide 0.956 carbonic acid 1.500

The specific gravity of carbon is undoubtedly much greater than any of the three. Now when oxygen gas combines with carbon, and forms carbonic acid, its specific gravity is increased, as happens in other cases. But how is it possible, says Berthollet, to conceive that the addition of carbon should diminish the specific gravity of oxygen gas, as would be the case if carbonic oxide were composed of these two ingredients only? Nay, carbonic acid, by dissolving an additional dose of carbon, would not only become specifically lighter than before, but even specifically lighter than oxygen gas, which Berthollet considers as incredible. But this reasoning is by no means sufficient to induce us to refuse credit to the conclusions of Cruikshanks; for similar instances are by no means so uncommon as Berthollet supposes. Mr Davy has given us an example which has some analogy to the case under our consideration.

The specific gravity of oxygen gas is 1.103

nitrous oxide 1.603
nitric oxide 1.094

Now when nitric oxide is converted into nitrons oxide by abstracting a portion of its oxygen, the heaviest of its component parts, its specific gravity is increased.

This is as great an anomaly as that at which Mr Berthollet startles.

^{*} Journals of the Royal Institution, i. 317.

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SECT. III.

OXIDES OF SULPHUR AND PHOSPHORUS.

ought now to proceed to the consideration of the of sulphur and phosphorus, which constitute the der of the oxide supporters; but these bodies are imperfectly known to admit of a separate dis-

The small number of facts which have been med were detailed in the first Book of this Work, the substances themselves, which constitute the these oxides, were under examination.

III. OXIDE SUPPORTERS.

cannot be formed by combustion, but make their cance in processes nearly the reverse of combustand hence it happens that the oxygen which they is still capable of supporting combustion. Cheave given the name of oxide to those oxides only base is azote; the oxide of muriatic acid has considered as an acid. This will prevent us from it into consideration in this Chapter; but a of some of its most important properties was when we were treating of its base.

Book II.

SECT. IV.

OF THE OXIDES OF AZOTE.

Azote and oxygen form two different oxides, both of which were discovered by Dr Priestley. They can only be exhibited in the state of a gas: Hence the first of them has been called nitrous oxide gas; the second, nitric oxide gas.

1. NITROUS ORIDE GAS.

History.

NITROUS oxide gas was discovered by Dr Priestley about the year 1776, and cailed by him deplicogisticated nitrous gas. The associated Dutch chen ists examined it in 1793, and demonstrated it to be a compound of azote and oxygen. But for a full investigation of its properties we are indebted to Mr Davy, who published an excellent dissertation on it in the year 1800. He gave it the name of nitrous oxide.

Prepara-

1. It may be procured by the following process: Take any quantity of nitrate of ammonia (a salt composed of nitric acid and ammonia) in crystals, and expose it in a retort, by means of a lamp, to a heat not under 340°, nor above 500°. It melts rapidly, and is decomposed, emitting a great quantity of gas, which is

Jour. de Phys. 2lii. 323.

[†] Researabes, chiefly concerning Natrous Cuide.

from the mouth of the retort, and may be received in jars in the usual manuer. The gas which over is nitrous oxide. This process was first out by Berthollet; but it was much simplified Davy .

Chap, &

Nitrous oxide gas, thus obtained, has all the me- Properties. properties of air: but it is much heavier than to specific gravity, according to Davy, is 1 603, Fair being 1.000. It is to common air nearly as 5 One hundred cubic inches of it, at the tempeof 60°, barometer at 30 inches, weigh 49.706

apable of supporting combustion even better mmon air; almost as well indeed as oxygen gas. We burns in it with a brillians flame and a crackvise. No combustible, however, burns in it, unbe previously brought to a state of ignition.

Priestley and the Dutch chemists had concluded cannot be respired; but they did not examine it thate of purity t. Mr Davy ascertained that it breathed for several minutes without any bad

The feelings produced by breathing it bear a resemblance to intoxication; but they are not ed by that langour and debility which is a con-

Proust has remarked, very jury, that the gas which comes the beginning of the process differs a little from nitrous oxide, It is lint mifrous gos.

By's have riber, p. 94.

Proceedey indeed found, in one instance, that a mouse breathed it fintes without oreasiness. It this experiment he seems to have it nearly pure. - Printiley, in 84.

Book 11. Division 11. longer than about four minutes, without the loss of voluntary motion altogether. When animals are confined in it, they give no signs of uneasiness for some ments; but they soon became restless, and, if not removed in a very few minutes, die altogether. Hence we see that, though this gas be respirable, it is much less so than common air or oxygen gas †.

Action of water.

3. This gaseous oxide is absorbed pretty rapidly by water, as Dr Priestley ascertained, especially when agitated. Water absorbs 0.86 parts of its bulk of this gas, or according to Dalton nearly its own bulk of it. It acquires a sweetish taste; but its other properties do not differ perceptibly from common water. The whole of the gas is expelled unaltered by boiling the water !.

Mr Davy describes the affects it had upon him as follows: "Having previously closed my nostrils and exhausted my lungs, I breshed four quarts of nitrous oxide from and into a silk bag. The first feeling were similar to those produced in the last experiment (giddines); but in less than half a minute, the respiration being continued, they diminided gradually, and were succeeded by sensations analogous to gentle pressure on all the muscles, attended by an highly pleasureable thrilling, particularly in the chest and the extremities. The objects around me became dazzling, and my hearing more acute. Towards the last impirations, the thrilling increased, the sense of muscular power became greater, and at last an irresistible propensity to action was indulged in; I recollect but indistinctly what followed; I know that my motions were various and violent.

[&]quot;These effects very soon ceased after respiration. In ten minutes i had recovered my natural state of mind. The thrilling in the extremities continued longer than the other sensations." Davy's Researches, p. 457. The gas has been breathed by a very great number of persons, and almost every one has observed the same things. On some few, indeed, it has no effects whatever, and on others the effects are always painful.

[†] Davy's Researches, p. 94.

[†] Priestley, ii. 8x.

When this gas combines with the water, it expels the common air which was formerly dissolved in the water. Hence the residuum of common air, which always appears when this gaseous oxide is exposed to a sufficient quantity of water ...

- 4. This gas is not altered by exposure to light, nor to any heat below ignition; but when made to pass through s red-hot porcelain tube, or when electric sparks are made to traverse this gas, it is decomposed, and converted into nitric acid and common air +.
- 5. There is no action between this gas and air or oxygen gas.
- 6. Sulphur, at the common temperature of the air, is Of the simnot altered by this gas. If it be introduced into it while tibles. burning with a blue flame, it is immediately extinguished; but if introduced while burning with a white flame, it continues to burn for some time with great brilliancy. and with a fine red flame. The products are sulphuric acid and azote. When about the half of the nitrous oxide is decomposed, the sulphur is extinguished I.

Phosphorus may be melted and sublimed in this gas without alteration; it may be even touched with a red hot wire without undergoing combustion; but when touched with a wire heated to whiteness, it burns, or rather detonates, with prodigious violence. The products are, azotic gas, phosphoric acid, and nitric acid: a part of the oxide remains undecomposed | .

Charcoal, confined in this gaseous oxide, may be kindled by means of a burning-glass. It continues to

[.] Davy. p. 89. Davy, ibid. p. 303.

burn with great brilliancy, till about the half of the gas is consumed. The products are carbonic acid gas and azotic gas *.

Hydrogen gas and nitrous oxide gas detonate violently with a red flame, when a strong red heat is applied, or when the electric spark is made to pass through the mixture. When the proportion of hydrogen is nearly equal to that of the oxide, the products are water and azote; when the proportion of hydrogen is small, nitric acid is also formed †.

Sulphureted, phosphureted, and carbureted hydrogen gas, likewise burn when mixed with nitrous oxide, and exposed to a strong red heat. The products differ according to the proportions of the gases mixed.

7. Neither azote nor muriatic acid appear to have any marked action on this gaseous oxide.

Of the me-

8. On some of the metals it acts with great energy at high temperatures. Thus iron wire burns in it with the same brilliancy as in oxygen gas, though the combustion lasts but a very short time. The iron is converted into black oxide; part of the nitrous oxide is decomposed, its azote is evolved, while its oxygen combines with the iron ‡. Zinc also may be oxidized in this gas §. Its effect upon the other metals has not been tried.

Azotites.

9. Oxide of azote is capable of combining with alkalies, and forming salts of a very peculiar nature; for the discovery of which we are indebted to the sagacity of Mr Davy. No combination takes place when the alka-

[#] Davy; p. 31 %

[†] Priestley, ii. 83. and Davy, p. 286.

^{\$ 1}bid. is. 86.

⁵ Davy, p. 117:

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we exposed to nitrous exide in the gaseous state. is it come into contact with them at the instant of rmation, it combines with them very readily. As combinations have not yet received a name, we e call them asotites till some better appeliation be ight of ".

westite of potash may be formed by the following Nitrous gas (a substance which will be descrilimmediately), by confining in it crystallized sulof potash t, is gradually deprived of a portion of beygen, and converted into nitrous oxide. If very y pulverised sulphite of potash, mixed with potash, exposed for a great length of time in a sufficient maity of nitrous gas, it is changed almost completeto sulphate of potash, while the oxide of azote, as it evolved, combines with the pure potash. Consemily the salt is converted into a mixture of sulphate totash and azotite of potash. The sulphate may be rated by solution, evaporation, and crystallization low temperature.

Azotite of potash is obtained in irregular crystals. composed of about three parts of alkali and one of nitrous oxide. It is soluble in water. Its taste mustic, and it has a peculiar pungency. It converts cable blues into green. Pulverised charcoal, mix-

Mr Davy has proposed to call them n tr mu, but this name is exmable, not only because it is contrary to the ideom of the English stage, but because it is inconsistent with the rules laid down for ing chemical terms.

Potash combined with sulphurous acid. This salt has a strong affifor oxygen. It absorbs it from nitrous gas, and is converted into her of parasts. Hence the change of natrous gas to nitrous oxide.

with it, and inflamed, burns with slight scintillations.

When projected into zinc in fusion, a slight inflammation takes place. All scids, even carbonic, seem capable of expelling the nitrous oxide from the potash the other properties of this salt have not been examined.

Azotite of soda may be formed in the same manner, and seems to agree nearly in its properties with azotite of potash. The nitrous oxide is disengaged from it by a heat of between 400° and 500°. Its taste is more serid than that of azotite of potash, and it seems to contain less oxide of azote †.

Mr Davy did not succeed in combining nitrous oxide with ammonia and earths; but he has rendered it probable that these azotites may be formed.

Composi-

ide gas just detailed, it is obvious that it is a supporter of combustion, and therefore that it contains oxygen in the same state as it exists in other supporters. That its other component part is azote, cannot be doubted, if we consider that either azote or nitric acid is constantly evolved when nitrous oxide is decomposed. The experiments of Mr Davy leave no doubt that these two substances are its only constituents. This philosopher found that 39 measures of nitrous oxide gas are capable of saturating 40 measures of hydrogen, and that after combustion the residue consists of 41 measures of azotic gas. But 40 measures of hydrogen were found to require 20.8 measures of oxygen ‡. From this it follows, that

† Ibid. p. s61.

Davy, p. 362.

if the component part of nitrous oxide were merely mixed together, the bulk of them would occupy a third more room than when combined; for 40 measures of nitrous oxide would be resolved into 20.8 measures of oxygen gas and 40 measures of azotic gas. But 20.8 cubic inches of oxygen gas weigh about seven grains, and 40 of azotic gas about 12 grains. Hence it follows, that nitrous oxide is composed of seven parts by weight of oxygen, and 12 of azote, or nearly

63 azote,

37 oxygen,

100

This statement coincides very nearly with the specific gravity of nitrous oxide. Thirty-nine cubic inches of nitrous gas, if the composition here given were precise, ought to weigh 19.38 grains. In reality they weigh 19.69 grains. The component parts of this gas, as obtained by the Dutch chemists, differ very little from the result of Mr Davy's experiments.

Much is still wanting to render the history of this singular substance complete. Mr Davy has laid open a very interesting field of investigation, which promises, if pursued far enough, to throw much light upon the nature of combustion: an operation more intimately connected with azote and its compounds than is at present supposed.

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I omit one measure, because the gases were not absolutely pure Vol. II.

II. NITRIC OXIDE.

History.

NITRIC OXIDE, usually denominated nitrous gas, was obtained accidentally by Dr Hales; but its nature and properties were investigated by Priestley, in one of the first excursions made by that illustrious philosopher into the then unbeaten tracts of pneumatic chemistry. As the phenomena exhibited by this oxide are intimately connected with the most important investigations in chemistry, its properties were examined with great care, and occupied the attention of almost every chemist of eminence.

Prepara-

1. It may be obtained by the following process: Put copper or mercury into a glass retort, and pour over it somewhat diluted nitric acid. The metal is rapidly dissolved with a strong effervescence, and a great quantity of gas issues from the mouth of the retort, which may be received in glass jars. This gas is nitrous gas.

Properties.

2. When pure it is invisible like common air, of which it possesses the mechanical properties. Its specific gravity is 1.094, that of air being 1.000, according to Davy *. Nitrous gas, then, is to common air nearly as 34 to 31. One hundred cubic inches of it, at the temperature of 60°, barometer 30 inches, weigh 33.923 grains.

Nitrous gas is exceedingly noxious to animals, producing instant suffocation whenever they attempt to breathe it.

^{*} Researches, p. 6.

The greater number of combustible bodies refuse to urn in it: A taper, for instance, is extinguished the Supports moment it is plunge into nitrous gas; the same thing oppens to sulphur, even though previously burning with a white flame. It is capable, however, of supporting the combustion of several bodies, as has been asceraned by the experiments of Priestley and Davy. When Homberg's pyrophorus is introduced into niyour gas, it takes fire spontaneously, just as it does in mmon air. Phosphorus, too, when introduced into his gas in a state of inflammation, burns with as much pleodour as in oxygen gas f.

4. When nitrous gas and common air are mixed toether, the mixture instantly assumes a yellow colour, eat is evolved, and the bulk of the two gases dimishes considerably; slowly, if the experiment be made wer mercury; but rapidly, if it be made over water. When the diminution has reached its maximum, the mature becomes perfectly transparent. The yellow blour is owing to a quantity of nitrous acid which is primed, and the diminution of bulk to the gradual abexption and condensation of this acid. What remains ther this absorption is only azotic gas. The cause of in remarkable phenomenon is obvious. The nitrous as combines with the oxygen of the air, and torms brous acid which is condensed; while the azote of the air remains behind in the form of a gas. Hence with equal quantities of nitrous gas and air the dimmuion of buth is always proportional to the quantity of

This substance will be described hereafter. The combustible part is charcoal and sulphur.

Davy, p. 134.

Book II. Division II. And ozygen. oxygen present in the air. Hence it informs us of the proportion of that substance in any particular air. The same phenomenon takes place when oxygen gas and nitrous gas are mixed; but the condensation is much more considerable. Indeed it would be complete, provided the two gases were perfectly pure, and mixed is the proper proportions.

From the experiments of Mr Dalton we learn that aitrous gas combines with two different proportions of oxygen gas. 21 measures of oxygen gas uniting either with 36 measures of nitrous gas, or with 72 measures.

When electric sparks are made to pass through nitrous gas, it is decomposed and converted into nitrous acid and azotic gas †.

Absorbed by water. an experiment of Mr Davy, it appears that 100 cubic inches of water at the common temperature, and proviously freed from air, absorb 11:8 cubic inches of nitrous gas, or nearly one-tenth, as Dr Priestley had ascertained; but the experiments of Dr Henry do not accord with this estimate. Water, by his trials, at the temperature of 60°, absorbs only five per cent. of its bulk of this gas 1. This solution has no particular taste, and does not redden blue vegetable colours. The gas is expelled again by boiling the water §; it separates likewise when the water is frozen ||.

Action of combustibles, 6. Nitrous gas is decomposed by phosphorus and charcoal at a very high temperature, and probably also

^{*} Phil. Mag. 22iii. 35 t.

^{\$} Flid Trans. 1803, p. 274.

[|] Priestley, il. 407.

Priostley, il. en.

⁵ Davy, p. 245.

by sulphur. These substances are converted into acids by combining with the oxygen of the gas while its azote ts evolved.

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Hydrogen gas mixed with it acquires the property of burning with a green flame. A mixture of these two gases does not take fire when electric sparks are made to pass through it; but according to Fourcroy, it deconstes when made to pass through a red hot porcelain tube; water is formed, and azotic gas is evolved *.

7. Nitrous gas has no action whatever on azotic gas, Incomboseven when assisted by heat. Neither does it act on muriatic acid.

8. Several of the metals have the property of decom- Andmetals. posing it, especially when assisted by heat. This is the case particularly with iron. Dr Priestley confined a portion of nitrous gas for some time in contact with a number of iron nails; the gas was converted into oxide of azote, in consequence, doubtless, of the iron abstructing part of its oxygen †. It was in this manner that nitrous oxide was discovered by that philosopher. When the iron is heated to redness by means of a burning-glass, the decomposition is complete, the whole of the oxygen is abstracted from the nitrous gas, and only azotic gas remains behind 1.

9. Dr Priestley ascertained that nitrous gas is absorbed by the green sulphate of iron; a property which is employed successfully to ascertain its purity. All that is necessary is, to expose a given portion of nitrous gas in a close vessel to the action of the green sulphate;

Fource y, ii, 91. 1 Iliid, p. 38.

the quantity of gas which remains unabsorbed gives the proportion of foreign bodies with which it is mixed. Mr Davy has proved, that all the salts containing the black oxide of iron possess the same property, and that they all absorb nitrous gas unaltered. The greatest part of it may even be expelled again by the application of heat. Several other metallic salts possess the same properties.

Converted into nitrous oxide.

10. The following bodies have the property of converting nitric oxide into nitrous oxide.

Alkaline sulphites,
Hydrogureted sulphurets,
Muriate of tin,
Sulphureted hydrogen gas,
Iron or zinc filings moistened with water.

To produce this effect, nothing more is necessary than to put these substances into jars filled with nitric oxide gas, and allow them to remain for a week or two. The substances gradually combine with a portion of oxygen, and are converted into oxides or salts †.

11. Nitrous gas is absorbed by alkaline solutions; but it does not appear from the experiments hitherto made, that it is capable, like oxide of azote, of combining with alkalies and earths, and forming salts.

Composi-

12. The conversion of nitric oxide gas into nitric acid, by combining it with oxygen, is a demonstration that it

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^{*} Davy, p. 179.

[†] Priestley and Davy, pessim.—During the action of the two last hadies on nitrous gas, ammonia is likewise formed.

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contains azote as a constituent part; and the property which several bodies have of absorbing oxygen from 11, and converting it into nitrous oxide gas, is a demonstration that oxygen is the only other ingredient which it contains. But it is by no means easy to ascertain the proportion of these two constituents. Mr Lavoisier, from a set of experiments made at an early period, and in a manner not susceptible of much accuracy, estimated their proportions at about 68 oxygen and 32 azote. But this estimate is irreconcileable with the experiments of other philosophers. The proportions obrained by Mr Davy deserve much more confidence, as his method was susceptible of greater accuracy. By means of a burning-glass he exposed to the action of the sun's rays a small portion of prepared charcoal, placed in a measured quantity of nitric oxide gas standing over mercury. The weight of the charcoal did not exceed a quarter of a grain, and the gas amounted to 16 very small measures. After the process was finished, the bulk of the gas was increased about 2ths of a measure. The whole of the nitric oxide was decomposed. Potash ley rapidly absorbed the whole of the gas except eight measures, which were pure azote. But the gas originally was found by experiment to contain 0.6 of a measure of azote. Therefore 15.4 measures of nitric oxide, when decomposed by charcoal, are converted into 16:15 measures; 7:4 of which are azote, and 8.75 carbonic acid. But 15.4 measures of nitric oxide weigh 5.2 grains, and 7.4 measures of azote weigh 2.2 grains. Hence it follows that 5.2 grains of nitric oxide contain 2.2 of azote; the remaining 3 grains must be oxygen. This gives us nitric oxide composed of 2.2 azote and 3 oxygen, or of 57.7 oxy-

gen and 42.3 of azote. The carbonic acid produced weighed 4.1 grains, and contained 1.15 grains of charcoal combined with 2.95 grains of oxygen, which it had absorbed from the nitric oxide. This gives us 5.2 grains of nitric oxide, composed of 2.95 oxygen and 2.25 azote; or 100 parts contain 57 oxygen and 43 azote, which differs very little from the last estimate. We may consider the last numbers as the nearest approximations to the composition of nitric oxide.

The proportion of oxygen contained in the compounds of azote, according to these results, may therefore be stated thus:

Azote.	Oxygen.	
100	58.7	Nitrous oxide
100	136	Nitric oxide
100	236	Nitric acid

Hence it follows that

*Davy, p. 129.

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CHAP. II.

OF ACIDS.

HEE word ACID, originally synonymous with Sour. ad applied only to bodies distinguished by that taste, been gradually extended in its signification by chemists, and now comprehends under it all substances pasessed of the following properties:

1. When applied to the tongue, they excite that sen- Properties. tion which is called sour or acid.

- 2. They change the blue colours of vegetables to a d. The vegetable blues employed for this purpose e-generally infusion of litmus and syrup of violets or raddishes, which have obtained the name of reagents tests. If these colours have been previously converted to a green by alkalies, the acids restore them main.
 - 3. They unite with water in almost any proportion.
- 4. They combine with all the alkalies, and most of be metallic oxides and earths, and form with them bose compounds which are called salts.

It must be remarked, however, that every acid does

not possess all these properties; but all of them possess a sufficient number of them to distinguish them from other substances. And this is the only purpose which artificial definition is meant to answer.

The acids are by far the most important class of bodies in chemistry. It was by their means indeed, by studying their properties, and by employing them as instruments in the examination of other bodies, that men of science laid the foundation of chemistry, and brought it to that state in which we find it at present. The nature and composition of acids, therefore, became a very important point of discussion, and occupied the attention of the most eminent cultivators of the science.

Opinion concerning the acid principle.

Paracelsus believed that there was only one acid principle in nature which communicated taste and solubility to the bodies in which it was combined. Beccher embraced the same opinion; and added to it, that this acid principle was a compound of earth and water, which he considered as two elements. Stahl adopted the theory of Beccher, and endeavoured to prove that his acid principle is sulphuric acid; of which, according to him, all the other acids are mere compounds. But his proofs were only conjectures or vague experiments, from which nothing could be deduced. Nevertheless, his opinion. like every other which he advanced in chemistry, continued to have supporters for a long time, and was even countenanced by Macquer. At last its defects began to be perceived: Bergman and Scheele declared openly against it; and their discoveries, together with those of Lavoisier, demonstrated the falsehood of both parts of the theory, by showing that sulphuric acid does not

That in the other soids, and that it is not composed of ther and earth, but of sulphur and oxygen.

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The opinion, however, that acidity is owing to some tinciple common to all the salts, was not abandoned. Wallerius, Meyer, and Sage, had advanced different beories in succession about the nature of this prinaple; but as they were founded rather on conjecture and analogy than direct proof, they obtained but few advocates. At last Mr Lavoisier, by a number of ingenious and accurate experiments, proved that several combustible substances when united with oxygen form scids; that a great number of acids contain oxygen; and that when this principle is separated from them, they lose their acid properties. He concluded, therefore, that the acidifying principle is oxygen, and that mids are nothing else but combustible substances comlined with oxy, en, and differing from one another acor ding to the nature of the combustible base.

Supposed oxygen.

This conclusion, as far as regards the greater number of acids, is certainly true. All the simple combustions, except hydrogen, are convertible into acids; and the acids are composed of oxygen and the combustic body combined. This is the case also with four the actals. It must not, however, be admitted with-

- 1. When it is said that oxygen is the acidifying prinple, it is not meant surely to affirm that oxygen possses the properties of an acid, which would be conary to truth; all that can be meant is, that it enters a component part into acids, or that acids contain it an essential ingredient.
- 2. But, even in this sense, the assertion cannot be ad-

that oxygen is an essential ingredient in all acids. Mut riatic acid, for example, has not yet been proved to contain oxygen, and the same observation applies to some other substances universally admitted among acids.

3. When it is said that oxygen is the acidifying principle, it cannot be meant surely to affirm that the combination of oxygen with bodies produces in all cases an acid, or that whenever a body is combined with oxygen, the product is an acid; for the contrary is known to every chemist. Hydrogen, for instance, when combined with oxygen, forms not an acid, but water, and the greater number of metallic bodies form only oxides.

The recent discoveries of Mr Davy render this, if possible, still more obvious. For we now know that the alkalies contain oxygen, and that they are as much indebted to that principle for their alkaline qualities at the acids are to it for their acid qualities.

All that can be meant, then, when it is said that oxygen is the acidifying principle, is merely that it exists as a component part in the greater number of acids; and that many acids are formed by combustion, or by some equivalent process. The truth is, that the class of acids is altogether arbitrary; formed when the greater number of the bodies arranged under it were unknown, and before any precise notion of what ought to constitute the characteristic marks of an acid had been thought of. New bodies, when they were discovered, if they possessed any properties analagous to the known acids, were referred without scruple to the same class, how much soever they differed from them in other particulars. Hence we find, under the head of acids, bodies which have scarcely a single property in common except that of combining with alkalies and earths. Chap. II. What substances, for instance, can be more dissimilar ban sulphuric, prussic, and uric acids? Hence the difsculty of assigning the general characters of the class of acids, and the disputes which have arisen about the propriety of classing certain bodies among acids. If we lay it down as an axiom that oxygen is the acidifying principle, we must either include among acids a great number of bodies which have not the smallest resemblance to those substances which are at present reck oned acids, or exclude from the class several bodies which have the properties of acids in perfection. The dass of soids being perfectly arbitrary, there cannot be such a thing as an acidifying principle in the most extensive sense of the word.

Berthollet has lately proposed a definition of acid, which, though not unexceptionable, gives us, I think, more precise notions of this class of bodies than those formerly entertained. Acids, according to him, are bodies capable of combining with alkalies, and of neutralizing them, while, at the same time, they lose their seidity ". According to this notion, the capacity of neutralizing each other constitutes the essential characters both of acids and alkalies; and that acid which is capable of neutralizing the greatest quantity of alkali possesses the character of acidity in greatest perfection. Were this definition admitted, several bodies at present arranged smong acids would be excluded from that class. greatest objection to it is the existence of some substances which occasionally act the part both of acids and

^{*} Statique Chemigar, 1, 69.

Book IL Division IL alkalies, and ought therefore to belong to both sets of substances. But perhaps these bodies (the metallic oxides) do not rigidly neutralize alkalies, and therefore approach more closely to that class. Besides, we can scarcely expect perfect precision in our arbitrary classifications.

Division of acids.

As the class of acids comprehends under it a considerable number of bodies, the subdivision of them under subordinate heads will be attended with considerable advantage. Now, all the acids, like the oxides, are either products of combustion, or supporters of combustion, or combustible. And upon examination, it must appear evident that each of these classes possesses such characters of distinction as to warrant their being considered separately. I shall therefore divide the acids into the three following classes: 1. Products; 2. Supporters; 3. Combustibles. The acids belonging to the two first classes have only a single base; but those belonging to the third have usually two or more bases, and are sometimes destitute of oxygen.

CLASS I.

OF ACID PRODUCTS.

ALL the acids which belong to this class possess the following properties.

- 1. They may be formed by combustion. Of course Properties their base is a combustible substance.
 - 2. They are incombustible.
- 3. They cannot be decomposed by the action of the most violent heat which can be applied *.
- 4. They are all decomposible by the joint action of a combustible body and caloric.
- 5. Oxygen is an essential ingredient in all of them. To them, therefore, the theory of Lavoisier applies with precision.

Some of the simple combustibles are capable of combining with two doses of oxygen, and forming acids with each. When that happens, the acid containing the minimum of oxygen is distinguished by the termi-

^{*} To this property there are some exceptions. It appears from the tate experiments of Gay-Lussac that sulphuric acid is decomposable by heat. Mem. d'Arcueil, i. 215.

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nation ous; that which contains the maximum, by the termination ic. Thus the acid of sulphur, with a minimum of oxygen, is sulphurous acid; the acid of sulphur, with a maximum of oxygen, is sulphuric acid. But it must be remarked, that this mode of distinguishing the doses of oxygen in acids is restricted to the first class of acids, and does not apply to the other two classes, except in one instance.

Names and composition. The acid products at present known are only five in number: but I am obliged to add to them, from analogy, two other acids, which have not hitherto been decompounded, but whose properties bear a much close resemblance to the products than to any of the other classes of acids. The following TABLE exhibits the names and constituent parts of these acids as far as they have been ascertained.

Names.	Bases.	Proportion of oxygen to 1900 base.
Sulphuric Sulphurous	Sulphur	136·5 88·6
Phosphoric Phosphorous	Phosphorus	114'7
Carbonic	Carbon	257
Fluoric Boracic	Unknown	

From this Table it is obvious that the acids belonging to this class are named from their bases. The two last, the component parts of which are unknown, are denominated from the substances which contain them most abundantly.

Chap. 11.

SECT. I.

OF SULPHURIC ACID.

THOUGH the ancients were acquainted with some of the compounds into which sulphuric acid enters, alum, or instance, and green vitriol, they appear to have been morant of the acid itself. It is first mentioned in the borks of Basil Valentine, which were published about the end of the 15th century. It was for a long time beained by distilling green vitriol, a salt composed of Alphuric acid and black oxide of iron; hence it was "Lled oil of vitriol, and afterwards vitriolic acid. Anther method of obtaining it was by burning sulphur oder a glass bell; hence it was called also oleum sul-Meris per campanam. The French chemists, in 1787, when they formed the new chemical nomenclature, gave the name of sulphuric acid.

1. At present it is generally procured by burning a Preparamixture of sulphur and nitre in chambers lined with had. The theory of this process is still somewhat obware. When sulphur is burnt in the open air or in oxgen gas, however high the temperature, only sulphuous acid is produced, and no appreciable quantity of sphuric acid. If the sulphur be mixed with black wide of manganese or any metallic oxide which readily erts with its oxygen, the combustion is more brilliant, at still sulphurous acid only is produced. No other Vol. 11.

method is known of producing sulphuric acid by the combustion of sulphur, excepting mixing it with nitrat Now the difficulty is to explain the pur which the nitre acts. The quantity used varies from a fifth to a tenth of the sulphur burnt; and according to the experiments of Chaptal, the best proportion is a seventh . Now the whole oxygen contained in this portion of nitre does not exceed the seventh part of the oxygen necessary to convert the sulphur into sulphuric acid. It is obvious, therefore, that the necessity of the nitre does not arise from its supplying the requisite quantity of oxygen. The most probable explanation has been furnished by Clement and Desormes. According to them, the nitric acid in the nitre is decomposed by the combustion, deprived of a portion of as oxygen, and emitted in the state of nitrous gas, while the sulphur is converted into sulphurous acid gas, which flies off mixed with the nitrous gas. The nitrous gas absorbs oxygen from the air of the apartment, and is converted into nitric acid, which immediately gives out in oxygen to the sulphurous acid and converts it into sulphuric acid. The nitrous gas is thus formed a second time; it absorbs oxygen a second time, and gives it away as before; and this process is repeated till the whole sulphurous acid is converted into sulphuric acid +.

The sulphuric acid, when first formed, is very weak, being diluted with the water necessary for condensing it; but it is made stronger by distilling off a portion of this water. By this process it is made quite trans-

Chimle appliquee aux arts. iii. 40.

Ann. de Com. lix. 329.

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ment; but itstill contains a little lead, which it dissolved am the vessels in which it was manufactured, and a litte potash, which it acquired from the nitre employed in swing the sulphur. To obtain it in a state of complete rity, the sulphuric acid of commerce must be distilled. This is easily done by putting it into a small retort with long beak. The bottom of the retort is placed uparting; while its beak is plunged half way into a ceiver, whose mouth it fits nearly, but not exactly. The said soon boils, and is gradually condensed in the eceiver. Too great a quantity should not be distilled tonce, otherwise the retort generally breaks in consequence of the violent agitation into which the boiling

g. Sulphuric acid is a liquid somewhat of an oily conistence, transparent and colourless as water, without
my smell, and of a very strong acid taste. When apisted to animal or vegetable substances, it very soon
testroys their texture.

and is thrown.

It always contains a quantity of water; part of which, owever, may be driven off by the application of a motrate heat. This is called concentrating the acid. Then as much concentrated as possible, its specific gratry is said to be 2.000; but it can seldom be obtained user than 1.85.

It changes all vegetable blues to a red except indigo. Coording to Erxleben, it boils at 546°; according to rgman, at 540°.

When exposed to a sufficient degree of cold, it crys-

Bergman, il. 152.

tallizes or freezes; and after this has once taken place, it freezes again by the application of a much inferior cold *. Morveau froze it at -4°; it assumed the ap, pearance of frozen snow. After the process began, it went on in a cold not nearly so intense. The acid melted slowly at 27.5°; but it froze again at the same temperature, and took five days to melt in the temperature of 43° ‡. Chaptal, who manufactured this acid, cace observed a large glass vessel full of it crystallized at the temperature of 45°. These crystals were in groups, and consisted of flat hexahedral prisms, terminated by a six-sided pyramid. They felt hotter than the surrounding bodies, and melted on being handled f. Chaptal has observed, that sulphuric acid, in order to erystallize, must not be too concentrated. This observation has been extended a good deal further by Mr Keir. He found that sulphuric acid, of the specific gravity of 1.780, froze at 45°; but if it was either much more or much less concentrated, it required a much greater cold for congelation ||. When as concentrated as possible, P find that it may be cooled down in thermometer tubes to the temperature of -36° before it congeals ¶.

Action of water.

3. Sulphuric acid has a very strong attraction for water. Neuman found, that when exposed to the atmosphere it attracted 6.25 times its own weight. Mr Gould found, that 180 grains of acid, when exposed to the atmosphere, attracted 68 grains of water the first day, 58 the second, 39 the third, 23 the fourth, 18 the

^{*} The freezing point was ascertained by the Duc d'Ayen in 1776. See Macquer's Dictionary.

^{\$} Encycl. Method. Chim. i. 376.

f. Phil. Trons. lxxvii. Part ii.

[§] Jour. de Phys. Expi. 473.

⁹ See Vol. I. p. 523.

hith, and at last only 5, 4, 3, 4, 3, 8cc. The 28th day Chap. 11. the augmentation was only half a grain *. The affinity therefore between sulphuric acid and water, as is the case in general with other substances, becomes weaker the nearer it approaches to saturation. He does not specify the specific gravity of his acid; but as it only attracted 3.106 times its own weight, it could not have been very concentrated.

The affinity between sulphuric acid and water becomes still more striking when these bodies are mixed together. If four parts of sulphuric acid and one part of ice, both at the temperature of 32°, be mixed together, the ice melts instantly, and the temperature of the mixture becomes 2120. On the other hand, if four parts of ice and one part of scid, at 92°, be mixed together, the temperature sinks to about -4°. The cause of this change of temperature has been already explained. When four parts of acid and one part of water are mixed together, the temperature of the mixture rises to about 8000 +. The density of this compound of acid and water is much greater than the mean. Heat is also evolved when other proportions of acid and water are mixed together, though not in so great quantity. Lavoisier and De la Place found, that when 2.625lbs. troy of sulphuric acid, of the specific gravity 1.87058, were mixed with 1.969lb. troy of water, as much caloric was evolved as melted 4.1226 lbs. troy of ice, or as much caloric as the acid and water would have given out had they been heated without mixture to 155.90 1. We have no reason to suppose that sulphuric acid, at

† Fourcroy, il. 65.

Phil. Trans. 1684.

³ Men. Par. 2780

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the density of 2.000, is free from all mixture of water: so far from that, we know for certain that it contains a considerable proportion; for when it is combined with other bodies (barytes, for instance, or potash), there is a considerable quantity of water which remains behind, and does not enter into the combination. Now, is it possible to determine how much real acid and how much water are contained in a given quantity of acid of a given specific gravity? Attempts were made to answer this important question by Homberg, by Bergman, by Wenzel, and by Wiegleb; but the differences between their determinations were too great to place any confidence in their accuracy. The subject has lately been resumed, and prosecuted with much industry and skill by Mr Kirwan, whose results may be considered as approaching as nearly to perfect accuracy as the present state of the science admits. His method was as follows: He dissolved 80 grains of potash in water, and saturated it exactly with sulphuric acid of a given specific gravity (we shall suppose of 2.00), and diluted it with water till its specific gravity was 1.019. The whole weight amounted to 3694 grains. Forty-five grains of sulphate of potash, dissolved in 1017 grains of distilled water, have the same specific gravity at the same temperature; whence it follows, that the proportion of salt in each was equal. But in the last solution the quantity of salt was 22.6 of the whole; therefore the quantity of salt in the first was $\frac{3694}{22.6} = 159.52$ grains. Now of this weight 86 grains were alkali; the remainder, therefore, which amounts to 73:52 grains, must be acid. But the quantity of acid employed was

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grains; of this there were 6.52 grains which did not ther into the combination, and which must have been ere water: 79 parts of acid, of the specific gravity 2, herefore, contain at least 6.52 parts of water; and esequently 100 parts of it contain 8'25 parts of wa-. It only remains now to consider how much water phate of potash contains. Mr Kirwan thinks it parains none, because it loses no weight in any debee of heat below ignition; and even when exposed a red heat for half an hour it hardly loses a grain. This is certainly sufficient to prove, at least, that is entains very little water; and consequently we may coclude, with Mr Kirwan, that 100 parts of sulhuric acid, of the specific gravity 2:000, are comlosed pretty nearly of 91.75 of pure acid and 8.25 of eter.

Since there is such a strong affinity between sulphuse acid and water, and since the density of the mixture different from the mean density of the ingredients, it becomes a problem of the greatest importance to determine how much of the strongest sulphuric acid that can prepared exists in any given quantity of sulphuric acid of inferior specific gravity, and which consequently consists of a determinate quantity of this strong acid duted with water.

This problem has been solved by Mr Kirwan. He mok sulphuric acid of the specific gravity 2.000, which the strongest that can be procured, for his standard, and the point was to determine how much of this stand-

^{*} Irisb Trans. iv. 18.

Book 11. Division 11. ard acid existed in a given quantity of acid of inferior density.

He concluded, from a number of experiments with sulphuric acid, of the specific gravities 1.8846, 1.8689, 1.8042, 1.7500 (for he could not procure an acid of the specific gravity 2.000 at the temperature of 60°, in which his experiments were performed), that when equal parts of standard acid and water are mixed, the density is increased by \$\frac{1}{2}\$th part of the whole mixture. Then, by applying a formula given by Mr Poujet*, he calculated, that the increase of density, on mixing

Mr Projet undertook the examination of the specific gravity of slephol mised with different quantities of water. He took for his send-and alcohol whose specific gravity was 8109, at the temperature of 65.75°. He then formed ten mixtures; the first containing nine measures of alcohol and one of water, the second eight measures of alcohol and two of water, and so on till the last contained only one measure of alcohol and nine of water. He took care that each of these measure should contain equal bulks, which he ascertained by weight, observing that a measure of water was to a measure of alcohol as 1 to 0 8 99. Thus tooks grains of water and \$199 of alcohol formed a mixture containing equal bulks of each. From the specific gravity of each of these mixture he discovered how much they had diminished in bulk in consequence of injurior, by the following method:

Calling A the real specific gravity of any of the mixtures; B its specific gravity found by calculation, supposing no diminution of bulk, a the number of measures composing the whole man; n-n the number to which it is reduced in consequence of mutual penetration—it is evident, since the increase of denuty does not diminish the weight of the whole mass, that $n = \frac{1}{2} \frac{1}{2$

The following Table contains the result of Mr Poujet's experiment, releulated according to that formula; the whole volume or a being = t

different quantities of standard acid and water, was as in the following TABLE:

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Measures of		Diminution of the whole vo- lume == 1 by	By calculation	
Water.	Alcohol.	experiment		
1	9	0.0100	0°0 03	
3	8	0.0187	0'0184	
3	7	010343	0.0242	
4	6	o-cz68	0.0276	
5	5	0.0788		
6		0.0166	0.0276	
7	3	0'0207	0'0242	
8	2	00123	0'0184	
9	*	0.0044	00103	

te is evident, from this Table, that the diminution of the bulk of the mixture follows a regular progression. It is greatest when the measures water and alcohol are equal, and diminishes as it approaches both ends of the series. Mr Poujet accounts for this by conceiving the alcohol to be dissolved in the water, which retains a part of it in its pures, or absorbe it. The quantity absorbed ought to be in the ratio of that of the solvent and of the body dissolved, and each measure of water will retain a quangats of alcohol proportional to the number of measures of alcohol in the mixture. Thus in a mixture formed of nine measures of alcohol and one of water, the water will contain a quantity of alcohol = 9; in one of eight measures of alcohol and two of water, the water will contain a quantity of alcohol = 8. Therefore the diminution of bulk in each mixwere is in a ratio compounded of the measures of alcohol and water which form it, in the Table given above, as 1 × 9, 2 × 8, 3 × 7, 4 × 6. &c. And in general, taking the diminution of both when the measures of both liquids are equal for a constant quantity, and calling it c, calling the number of measures s, the number of measures of alcohol s, the in-

Number of Number of parts of stan dard acid.		Augmenta- tion of den- sity.	
5	95	0.0252	
10	90	0.0479	
15	85	0.0679	
20	80	0.0856	
25	75	0.0555	
30	70	0.1119	
35	65	0.1213	
40	60	0.1279	
45	55	0.1319	
50	50	0.1333	

trease of density or diminution of bulk π ; we shall have $\epsilon: \pi: \frac{\pi}{2} \times \frac{\pi}{2}$: $\pi - \pi \times \pi$ and $\pi = \frac{4\epsilon}{\pi^2} \times \pi \times \pi^2$, or (making $\pi - \pi$) $- 4\epsilon = -4\epsilon \pi^2$.

The diminution of bulk, calculated according to this formula, makes the last column of the Table in this note. They correspond very well with experiment, while the measures of alcohol are more than these of water, but not when the reverse is the case. This Mr Poujet thinks is owing to the attraction which exists between the particles of water, and which, when the water is considerable compared with the alcohol, resists the union of the water with the alcohol.

By the formula $x = \frac{4 c \pi \kappa - 4 c \kappa^2}{\kappa^2}$, the quantity of alcohol of the standard may be determined in any mixture where the alcohol exceeds the water.

By adding these augmentations to the specific gravity Chap II. of the above mixture, found by calculation, and taking the mean for the intermediate quantities, he drew up a table of the quantity of acid of the density 2.00, contained in a given weight of acid of inferior density, from 2 to 1'4666. Sulphuric acid of this last density contains just 0.5 of sulphuric acid of 2. The quantity contained in acids of inferior densities was ascertained from actual observation. He found by the first part of the table, that 100 parts of acid, of the specific gravity 1.8472, contained \$8.5 parts standard; consequently 400 grains of this acid contain 354 grains standard. He took six portions of this acid, each containing 400 grains, and added to them as much water as made them contain respectively 48, 46, 44, 42, 40, 38 grains standard. The quantity of water to be added, in order to produce this effect, he found by the following method: Suppose x = the quantity of water to be added to 400 parts of acid, that the mixture may contain 48 per cent. of standard acid. Then 400 + x : 354 : 100 : 48, and consequently x = 337.5. After finding the specific gravity of these, the half of each was taken out, and as

Whole,
$$1 - x \times a + b \times = 1 - 4c \times + 4c x^{2} \times y$$
.

Hence $x = cos - \frac{a - b}{8cy} + \sqrt{\frac{a - b}{4cy} + (\frac{a - b}{8cy} - cos)}$
 $y = \frac{a - a + b \times a}{1 - 4c \times + 4c \times^{2}}$

And making $x = cos + cos = c$

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much water added; and thus the specific gravities corresponding to 24, 23, 22, 21, 20, 19, were found. Then six more portions, of 400 grains each, were taken, of the specific gravity 1-8393, and the proper quantity of water added to make them contain 36, 34, 32, 30, 28, 26 per cent. of standard. Their specific gravities were found, the half of them taken out, and as much water added; and thus the specific gravity of 18, 17, 16, 15, 14, and 13 found. Care was taken, after every addition of water, to allow the ingradients sufficient time to unite.

By multiplying the standard acid, as ascertained in this Table, by 0.8926, he obtained the quantity of real acid contained in sulphuric acid of different densities at the temperature of 60°. This may be seen in the following TABLE*.

Strength of sulphuric acid.

100 Parts Sp. Gravity.	Real Ackl.	100 Parts Sp. Gravity.	Real Acid.
2.0000	89.29	1.8070	75.80
1.9859	88.39	1.7959	75.—
1.9719	87.50	1.7849	74.11
1.9579	86.61	1.7738	73.28
1.9439	85.71	1.7629	72.32
1.9299	84.82	1.7519	71.43
1.9168	83.33	1.7416	70.54十
1.9041	63.04	1.7312	69.64
1.8914	82.14	1.7208	68.75
1-8787	81.25	1.7104	67.86
1.8660	80.36	1.7000	66.98
1.8542	79.46	1.6899	66.07
1.8424	78.57	1.0800	65.18
1.8306	77.68	1.6701	64.58
1.8188	76.79+	1.6602	63.30

^{*} Nicholson's Journal, iii. 213.

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	oo Parte c. Gravity.	Real Acid.	soo Parts Sp Gravity.	Roal Acid
Π	1.0503	62.20	1.2951	31.25
L	1.6407	61.61	1.2847	30*35
u	1.6312	60.71	1.2757	29.46
П	1.0217	59.82	1.2668	28.57
u	1.6122	58.93	1.2589	27.68
П	1.6027	58.03	1-2510	26.78
H	1.5932	57.14	1'2415	25.89
н	1.5840	56.25	1.2320	25'
н	1-5748	55.36-	1.2210	24.10
	1-5656	54.46	1.5101	23.21
	1.5564	53-57	1.2009	22.32
u	1.5473	52.68	1.1918	21.43
П	1.5385	51.78	1.1836	20.53+
п	1.5292	50.89	1.1745	19'64
Ł	1.5202	50100	1.1018	18-75
L	1.2112	49.11+	1.1614	17.85
ı	1.2022	48.21	1.1231	16.92
L	1.4933	47.32	1.1358	16.01
1	1.4844	46'43	1.1308	15'18-
П	1.4755	45*53	1.1308	14.28
Ш	1.4566	44.04	1.1159	13.39
н	1.4421	43.75	1.1011	12.50
ł	1.4189	42'86+	1.0955	11.60
I	1.4099	41.96		
п	1.4010	41.07	1.0896	10.71
Н	1.3875	40-18	1.0833	9.80
П	1.3768	30.58	1.0780	8193-
F	1.3663	38*39	1.0725	8.03
	1.3286	37.50	1.0606	7'14
1	1.3473	35.60	1.0610	6.52
1	1.3360	35.41	1.0555	5.35
	1.3254	34.82	1.0492	4-40
ł	1.3149	33.83	1.0450	3.2.
	1.3105	33.03	1.0396	2.61
I	1:3056	32.14	1.0343	1.78

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Its composition.

4. Mr Lavoisier attempted to ascertain the proportion of the constituents of this acid, by measuring the quantity of oxygen absorbed by a given weight of sulphur during its combustion. His result was 71 parts of sulphur and 29 of oxygen. But this method was not susceptible of sufficient precision to warrant much confidence. Mr Thenard had recourse to a much better method *, which was employed still more lately for the same purpose by Mr Chenevix with much address †. Nitric acid was distilled off 100 parts of pure sulphur repeatedly till the whole sulphur was converted into an acid. The sulphuric acid, thus formed, was separated by means of barytes, with which it forms an insoluble compound. The 100 parts of sulphur, thus acidified, yielded 694 parts of dry sulphate of barytes. 100 parts of sulphate of barytes contain 14.5 parts of sulphur. This experiment was repeated by Klaproth with the same result. To know the portion of oxygen in sulphuric acid, it is only necessary to know the weight of sulphuric acid contained in dry sulphate of barytes. A great many experiments have been made to determine that point. Here Mr Chenevix seems to have fallen into an error. He makes it only 24 per cent. The analysis of Thenard indeed nearly agrees with his; but all other chemists make it above 30 per cent. I consider the result obtained by Kirwan and Klaproth as the most accurate. They coincide, and find sulphate of barytes composed of 33 parts sulphuric acid and 67 of barytes. Hence it follows, that sulphuric acid

[#] Ann. de Chim. xxxii. 266. † Irish Trans. 2802, p. 233.

s composed of 14.5 of sulphur, and 18.5 of oxygen; or of

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42.3 sulphur 57'7 oxygen

100.0

or 100 sulphur and 136.5 oxygen .

5 Sulphuric acid is not altered by the action of light Action of nor caloric. It does not combine with oxygen. It was oxygen, affirmed indeed by some chemists, that sulphuric acid might be combined with oxygen by distilling it off the black oxide of manganese; but the assertion was refuted by the experiments of Vauquelin.

b. None of the simple combustibles act upon it at the Of the sime usual temperature of the atmosphere, or at least their tibles. action is so slow as not to be perceptible. But when they are assisted by heat, they are all capable of deposing it.

When sulphur is boiled in this acid, it absorbs a portion of its oxygen, or at least combines with it, and the whole is converted into sulphurous acid. Phosphorus also absorbs oxygen from it by the assistance of heat. sulphurous acid is driven off, and phosphoric acid formed. At the boiling temperature charcoal also absorbs oxygen from it, and converts it into sulphurous acid. At a red heat it even converts it into sulphur. When hydrogen gas and sulphuric acid are made to pass together through a red hot tube of porcelain, the acid is completely decomposed, water is formed, and sulphur deposited †.

Book II. Division II. Of the incombunibles, 7. Azote has no action on sulphuric acid; but this acid readily absorbs muriatic acid, and forms with it a liquid of a brownish tinge, which emits the dense and suffocating odour of muriatic acid, and corrodes vegetable and even metallic bodies near which it happens to be placed.

Of metals,

8. When zinc or iron is thrown into sulphuric acid, a violent action takes place, if the acid be diluted; water is decomposed, its hydrogen flies off, and its oxygen / combines with the metals. If the acid be concentrated, the action is much less violent, and sulphurous acid ex-Upon tin and copper the acid acts very slowly and feebly, unless its action be assisted by heat when it oxidizes and dissolves them. On silver, mercury, antimony, bismuth, arsenic, and tellurium, it does not act except at a pretty high temperature. These metals abstruct part of its oxygen, and convert one portion of it to sulphurous acid, while another portion combines with the oxides thus formed. When boiling hot it oxidizes lead, and dissolves cobalt, nickel, and molybdenum: But it has no perceptible action on gold nor platinum at any temperature to which it can be raised.

Of sikalice and earths.

- 10. It unites readily with all the alkalies and earths except silica, and with most of the metallic oxides, and forms salts denominated sulphates. Thus the combination of sulphuric acid and soda is called sulphate of soda; the compound of sulphuric acid and lime, sulphure of lime, and so on.
- 11. It absorbs a very considerable quantity of nitrous gas, and acquires by that means a purplish colour.

This is one of the most important of all the acids, ot only to the chemist but to the manufacturer also; eing employed to a very great extent in a variety of nanufactures, especially in dyeing.

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SECT. II.

OF SULPHUROUS ACID.

Though some of the properties of this acid must have History. been known in the remotest ages, as it is always formed during the slow combustion of sulphur, Stahl was the first chemist who examined it, and pointed out its peculiar nature. He gave it the name of phlogisticated rulphuric acid, from the supposition that it is a compound of sulphuric acid and phlogiston. His method of procuring it was to burn sulphur, and expose to its flames cloth dipped in a solution of potash. By this method he obtained a combination of potash and sulphurous acid; for sulphur forms by combustion only sulphurous acid. Scheele pointed out, in 1771, a method of procuring sulphurous acid in quantities *. On Stahl's salt he poured a portion of tartaric acid, and then applied a gentle heat. The sulphurous acid is in this nanner displaced, because its affinity for potash is not so strong as that of tartaric acid; and it comes over in-

^{*} Scheele, i. 43.

to the receiver dissolved in water. Dr Priestley, in 1774, obtained it in the gaseous form, and examined it properties while in a state of purity. Berthollet published a dissertation on its formation, composition, and uses, in 1782 and 1789; and in 1797 appeared a very complete dissertation on its combinations by Fourcey and Vauquelin ‡.

Prepara-

1. Sulphurous acid may be procured by the following process: Put into a glass retort two parts of miphuric acid and one part of mercury, and apply the heat of a lamp: The mixture effervesces, and a gas issues from the beak of the retort, and may be received in glass jars filled with mercury, and standing in a mercarial trough. This gas is sulphurous acid.

Properties.

- 2. Sulphurous acid, in the state of gas, is colourless and invisible like common air. It is incapable of maintaining combustion; nor can animals breathe it without death. It has a strong and suffocating odour, precisely the same with that exhaled by sulphur burning with a blue flame: sulphur, by such a combustion, being totally converted into sulphurous acid. Its specific gravity, according to the experiments of Kirwan, is 2,265, that of air being 1,000: At the temperature of 60°, barometer 30 inches, 100 cubic inches of this gas weigh 70,215 grains.
- 3. This acid reddens vegetable blues, and gradually destroys the greater number of them. It exercises this power on a great variety of vegetable and animal colours. Hence the use of the sumes of sulphur in bleach-

[#] On A'r, ii. 295. † Mem. Par. 1782. Ann. de Chim. ii. 54. ‡ Ann. de Chim. Exiv. 229. | Kirwan on Phlogiston, p. 30.

me wool and in whitening linen steined by means of Chap. It. firmits.

Planche has observed, that when syrup of violets, previously reddened by acids, is mixed with a little sulphurous acid, the original blue colour of the symp is restored, and its intensity gradually dintinishes ".

4. Dr Priestley discovered, that when a strong heat applied to this acid in clove vessels, a quantity of sulphur is precipitated, and the acid is converted into sulphuric +. Berthollet obtained the same result; but Foureroy and Vauquelin could not succeed 1.

According to Clouet and Monge, when this gas, in a state of condensation, is exposed to the temperature of -18°, it is condensed into a liquid f.

5. Water absorbs this acid with rapidity. According Action of to Dr Priestley, 1000 grains of water, at the temperature 54 5", absorb 39 to grains of this acid. Foureroy. on the other hand, affirms, that water at 40° absorbs the third of its weight of sulphurous acid gas ¶. Upon trying the experiment, I found that one cubic inch of water at the temperature of 61°, barometer 29.55 inches, absorbs 33 inches of gas. But an inch of water weighs about 25311 75 grains, and 39 inches of this gas weigh 25-16. Hence it follows, that 1000 water in weight absorbs \$1.5 of this gas. This exceeds considerably the estimate of Priestley, though it comes short of that of Fourcroy. Water impregnated with this gas acquires, according to my experiments, the specific graviby 1.0513, even in as high a temperature as 68°. This

1 Nicholann's Journal, i. 313

¶ Ibid. p. 77.

Ann. de Gbim. Ix. 253.

⁴ On Air, in 330.

Fourtruy, il. 74.

water may be frozen without parting with any of the acid gas. When water, which has been saturated with this acid at the freezing temperature, is exposed to the heat of 65.25°, it is filled with a vast number of bubbles, which continually increase and rise to the surface. These bubbles are a part of the acid separating from it. It freezes a few degrees below 32°*.

Of oxygen and air,

spheric air or to oxygen gas, it gradually combines with oxygen, and is converted into sulphuric acid. This change takes place more completely if the acid be combined with an alkali or earth. When a mixture of sulphurous acid gas and oxygen gas is made to pass through a red hot porcelain tube, the two bodies combine, and sulphuric acid is formed †.

Of the other simple bodies,

- 7. Of the simple combustibles, sulphur and phosphorus have no action on it whatever; hydrogen gas and charcoal do not alter it while cold, but at a red heat they decompose it completely: water or carbonic acid is formed, and sulphur deposited ‡.
- 8. Neither azote nor muriatic acid produce any change on it.
- 9. Sulphurous acid does not seem capable of oxidizing or dissolving any of the metals except iron, zinc, and manganese.
- 10. It combines with alkalies, earths, and metallic oxides, and forms salts known by the name of sulphites.

11. Sulphuric acid absorbs this gas in considerable

Of sulphuric acid.

^{*} Fourcroy and Vauquelin, Nicholson's Jour. i. 313

[†] Priestley, ii. 332. Fourcroy, ii. 74.

[†] Fourcroy and Vauquelin.

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entity. It acquires a yellowish brown colour, a petrating odour, and the property of smoking when exmed to the air. When this mixture is distilled, the est vapour which comes over, and which is a comand of the two acids, crystallizes in long white isms. This singular compound, formerly known by name of glacial sulphuric acid, smokes in the air ; when the atmosphere is moist, melts with effervesnce. When thrown into water, it hisses like a red iron. It has the odour of sulphurous acid . This prious compound, first mentioned by Kunkel, and afrwards by Bohn and by Neuman, attracted a great al of the attention of chemists, who attempted to inestigate the cause of its singular properties. The geeral opinion, in consequence of the experiments of Lellot, Wiegel, Meyer, and Gottling, was, that phloiston, some how or other, contributed to its produc-Dolfusz + made a set of experiments on it in 1784; from which he concluded that it was peculiar sulphate of iron, by the distillation of which it had en usually procured. These experiments were recated and varied by Morveau 1; and this philosopher onsidered it as probable that the glacial acid is merely alphuric acid, totally deprived of moisture. y has lately demonstrated that it is a compound of aphuric and sulphurous acids.

12. As this acid is formed by the combustion of sultur, it cannot be doubted that it is composed of the time ingredients with sulphuric acid; and as it is evol-

Composi-

[•] Fourcroy, ii. 78. † Crell's Annale, 2785, i. 438.
5 Sorye, Method. Chim. i. 390.

ved from sulphuric acid by the action of sulphur, and likewise by some of the metals, it cannot be doubted that it contains a smaller proportion of oxygen. I endesvoured to estimate the constituents of this acid in the following manner: By causing a current of sulpharous acid gas to pass through a solution of carbonate of potash in water, I formed the saft called sulphite of petash. When this salt, previously dried upon bleating paper, is exposed to the heat of 300°, it loses 313 pm cent, of its weight. When 100 grains of it are heated to redness in a platinum crucible, the sait decrepitates, becomes of an opaque white, and emits a hine same. By this treatment it loses 22.3 per cent. and leaves as a residunm sulphate of potash. This sulphate of potash contains 22.25 of sulphwric soid, the rest is potash, From this analysis it follows that sulphite of potach is composed of

> 22.30 voiatile matter 22.25 sulphuric acid 55.45 potash

100:00

When 100 grains of sulphite of potash are heated in a retort attached to a mercurial apparatus, there come over 18 cubic inches of sulphurous acid gas, 5'1 grains of sulphur, and a little water. The remaining salt weighs 77'7 grains, and contains 23'2 sulphuric acid, the rest potash. From this analysis it follows that the 100 grains of sulphite of potash were resolved into

SULPHUROUS.

29.2 sulphuric acid
5.1 sulphur
54.5 potash
17.2 sulphurous acid and water

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100.0

23.2 sulphuric scid and 5.1 of sulphur must en united in the salt, and formed sulphurous 2f course, sulphurous acid is composed of 23.2 ic acid and 5.1 sulphur, or, per cent. of

82 sulphuric acid

18 sulphur

we know the constituents of sulphuric acid, it is nee to deduce the following as the proportion agredients of sulphurous acid:

> 53 sulphuc 47 oxygen

100

sulphur and 88.6 oxygen.

SECT. III.

OF PHOSPHORIĆ ACID.

HORUS, like sulphur, forms an acid with two proportions of oxygen. Combined with a m of oxygen, it constitutes phosphoric acid;

with a smaller proportion, it constitutes phosphorous

Phosphoric acid remained unknown till after the discovery of phosphorus. Boyle is perhaps the first chemist who mentions it; but Margraff first examined its properties, and demonstrated it to be a peculiar acid. Its properties were afterwards more completely investigated by Bergman, Scheele, Lavoisier, Pearson, Fourcroy, and Vauquelin, and several other distinguished chemists.

Prepara-

1. It may be formed by setting fire to a quantity of phosphorus contained in a wessel filled with oxygen gas. The phosphorus burns with great rapidity, and a great number of white flakes are deposited, which are plosphoric acid in a state of purity. It may be obtained too by heating phosphorus under water till it melt, and then causing a stream of oxygen gas to pass through it by means of a tube. In this case the acid as it forms combines with the water; but the liquid may be evaporated off by the application of heat, and then the acid remains behind in a state of purity. But the usual method of procuring it is to throw phosphorus in small pieces at a time into hot nitric acid. A violent effervescence takes place, the phosphorus combines with oxygen, and nitrous gas is emitted. After the whole of the phosphorus is acidified, the liquid is to be evaporated to dryness, in order to drive off the remains of nitric acid which may not have been decomposed. This process was first put in practice by Lavoisier. be taken not to apply too much heat, not to add too much phosphorus at once, and not to have the nitric acid too strong; otherwise the phosphorus takes fire, and usually breaks the vessels in pieces. When the acid bydrogen gas, which burns and occasions a considerable loss. This loss may be diminished by cautiously dropping nutric acid into the hot mass.

2. The acid thus prepared may be put into a platinum crucible, and heated to redness, to drive off all the
water. It is then in a state of purity. It is solid, colourless, and transparent, and not unlike glass in appearance. It reddens vegetable blues; it has no smell;
its taste is very acid, but it does not destroy the texture
of organic bodies.

When exposed to the open air, it soon attracts moisture, and deliquesces into a thick oily-like liquid; in which state it is usually kept by chemists. When exposed to the fire in a platinum crucible, its water gradually evaporates, and leaves it in the state of a transparent jelly. If the heat be increased it boils and bubbles up, owing to the separation of the remainder of its water, accompanied with a small portion of acid. At a red heat it assumes the form of a transparent liquid, and when cooled resembles the purest crystat. In this state it is known by the name of phosphoric glass. This glass is merely phosphoric acid totally deprived of water. It has an acid taste, is soluble in water, and deliquesces when exposed to the air. At a red heat it evaporates very copiously in a white smoke, which has no strong odour, unless the evaporation be prevented by the presence of a base.

The specific gravity of this acid, in a state of dryness,

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is 2.687°; in the state of glass, 2.8516†; in the state of deliquescence, 1.417‡.

Action of water.

3. This acid is very soluble in water. When in the state of white flakes, it dissolves with a hissing noise, similar to that made by red hot iron plunged into water. When in the state of glass it dissolves much more slowly. The heat evolved during the combination of this acid and water is much inferior to that evolved when sulphuric acid enters into a similar combination. Phosphoric acid obtained by deliquescence, when this ed with an equal quantity of distilled water, at quired so little beat as to raise the thermometer only one degree, as Mr Sage observed. Mr Lavoisfer raised the thermometer from 50° to 63° by mixing phosphoric acid, boiled to the consistence of a syrup, with an equal quantity of water; and from 50° to 104° when the acid was as thick as turpentine §.

Action of the combustibles,

- 4. Oxygen gas has no action on phospholic acid, whatever be the temperature. Neither is it detemposed or altered by any of the simple combustibles, if we except charcoal ¶; which, though it has no action on it while cold, at a ted heat decomposes it completely; cat-bonic acid is formed, and phosphorus sublimed. This is the common process for obtaining phosphorus.
- 5. None of the simple incombustibles have my particular action on it.
 - 6. This acid, when in a liquid state, is capable of oxi-

Of metals.

Bergman's Sciegrophie, p 20. Eng. Tress.

f Hamenfratz, Ann. de Chim. xxviii. 11.

^{\$} Encyc. Methe . Chim. i. 221.

⁵ Keir's Dictionery

And sethis also hydrogen; but the experiment has not been mada

Paing some of the metals, especially when assisted by beat; at the same time hydrogen gas is emitted. Hence we see that the oxidizement is owing to the decomposition of water. In this manner it oxidizes iron, tin, lead, zinc, antimony, bismuth, manganese; but on some of these it acts very slowly. When fused with several of these metals, as tin, iron, and zinc, it is converted into phosphorus; a proof that they have a stronger affinity for oxygen.

It does not act upon gold, platinum, silver, copper, mercury, arsenic, cobalt, nickel. It appears, however, to have some action on gold in the dry way, as it is called; for when fused with gold-leaf it assumes a purple colour: a proof that the gold has been oxidized.

7. Phosphoric acid combines with alkalies, earths, Phosphates. and metallic oxides, and forms with them salts known by the name of phosphates.

8. The component parts of this acid have been ascer- Compone tained in a more satisfactory manner than almost any tion. other chemical compound. Mr Lavoisier deduced from his experiments, that 45 parts of phosphorus, when burnt in oxygen, absorbed about 69-375 parts of that gas, and produced about 114 parts of phosphoric acid . But his mode of experimenting was not susceptible of much precision. It consisted in burning phosphorus in oxygen gas, and ascertaining the increase of weight, which he ascribed to oxygen. Mr Rose has lately published an elaborate set of experiments on the constituents of phosphoric acid. He acidified the phosphorus by means of nitrie acid, and ascertained the weight by combining

Lavoister's Elemente, p. 104.

the acid formed with oxide of lead. The result of his experiments was, that the acid was composed of

46.5 phosphorus
53.5 oxygen

0.00E

or 100 phosphorus combines with 114.75 of oxygen, and forms 214.75 of phosphoric acid. Some experiments, which I made in the same way nearly, gave me a result a good deal different. I combined the acid formed with lime. But as I do not recollect the proportion of acid which I estimated in phosphate of lime, it is impossible to say how far my results differed from those of Rose.

This acid is too expensive to be brought into common use. If it could be procured at a cheap rate, it might be employed with advantage, not only in several important chemical manufactures, but also in medicine, and perhaps even for the purposes of domestic economy.

SECT. IV.

OF PHOSPHOROUS ACID.

History.

THE acid obtained by the burning of phosphorus differs according to the rapidity of the combustion; or, which is the same thing, according to the temperature

[#] Gehlen's Jour. 2d. Series, ii. 309.

Chap. II.

in which the process is conducted. When heated to 148°, it burns rapidly, and the product is phosphoric acid; when allowed to burn gradually, at the common temperature of the air, the product is phosphorous acid, which contains a smaller proportion of oxygen. The difference between these two acids had been remarked by Sage, by Proust, and by Morveau; but it was Lavoisier who first, in 1777, demonstrated that they form different compounds with other bodies, and that the difference between them is owing to the different proportions of oxygen which they contain *.

- 1. Phosphorous acid is prepared by exposing phos- Preparaphorus during some weeks to the ordinary temperature of the atmosphere. Even in winter the phosphorus undergoes a slow combustion, and is gradually changed into a liquid acid. For this purpose, it is usual to put small pieces of phosphorus on the inclined side of a glass funnel, through which the liquor which is formed drops into the bottle placed to receive it. From one ounce of phosphorus about three ounces of acid liquor may be thus prepared. It was called phlogisticated phosphoric acid by Morveau, from the supposition that it is a compound of phosphoric acid and phlogiston.
- 2. Phosphorous acid, thus prepared, is a viscid liquid, Properties. of different degrees of consistence, adhering like oil to the sides of the glass vessel in which it is contained. emits the smell of garlic, especially when heated. taste is acid like that of phosphoric acid, and it produces the same effect upon vegetable colours. Its specific gravity has not been determined.

· Mem. Par. 1777.

3. It combines with water in every proportion; but it cannot, like phosphoric acid, be obtained in a concrete state.

Action of Water.

When heated, part of the water which it contains is at first evaporated. Then large bubbles of air rise to the surface; there they break, and emit a dense white smoke, or even take fire if the experiment be performed in an open vessel. The emission of these bubbles of phosphureted hydrogen gas continues for a long time. When the process is finished, the acid which remains is no longer phosphorous, but phosphoric acid. These phenomena show us, that phosphorous acid holds in solution a certain portion of phosphorus, which at a high temperature decomposes water, and is converted partly into phosphoric acid, partly into phosphureted hydrogen.

Of simple bodies.

- 4. This acid is converted into phosphoric acid by exposure to air or oxygen gas. The process is exceedingly slow, and the conversion is never complete. It succeeds better when the acid is diluted with a great proportion of water *.
- 5. Phosphorous acid is not acted upon by any of the simple combustibles except charcoal, and perhaps also hydrogen. Charcoal decomposes it at a red heat a well as phosphoric acid. The products are carbon acid and phosphorus. It does not act on the simple incombustibles.
- 6. Its action on metals is exactly similar to that phosphoric acid, excepting only that the hydrogen gavelved during the oxidizement of the metals have fetid smell, and holds phosphorus in solution.

^{*} Fourcroy, ii. 55.

1. It combines with alkalies, earths, and metallic oxa, and forms compounds distinguished by the name Phone Mosphites.

. Sulphuric acid produces no change upon it while Action of id; but at a boiling heat it parts with some of its ygen, and the phosphorous acid is converted into asphoric acid. Nitric acid also, when assisted by w, converts it readily into phosphoric acid. This mishes us with by far the best process for obtaining posphoric acid at present known. Mix phosphorous id, obtained by slow combustion, with one-eighth of weight of nitric acid of the specific gravity 1.3, and The natric acid is decomposed, and pure phose poric acid remains behind. For this process we are edebted to Fourcroy ..

This acid has not hitherto been put to any use. The istory of its preparation is sufficient to convince us, hat it is composed of the same constituents as phosphois acid. But the exact proportion of these constituents not hitherto been ascertained.

SECT. V.

OF CARBONIC ACID.

the rapid progress of chemistry, during the latter History of the 18th century, was in a great measure owing

[·] Fources, il. 86.

to the discovery of this acid, it may be worth while to trace the history of it somewhat particularly.

Paracelsus and Van Helmont were acquainted with the fact, that air is extricated from limestone during certain processes, and the latter gave to air thus produced the name of gas. Hales ascertained the quantity af air that could be extricated from these bodies in different processes, and showed that it formed an essential part of their composition. Dr Black proved that the substances then called lime, magnesia, and alkalies, are compounds, consisting of a peculiar species of air, and pure lime, magnesia, and alkali. To this species of air he gave the name of fixed air, because it existed in these bodies in a fixed state. This air or gas was afterwards investigated by Dr Priestley, and a great number of its properties ascertained. From these properties Mr Keir* first concluded that it was an acid; and this opinion was soon confirmed by the experiments of Bergman, Fontana, &c. Dr Priestley at first suspected that this acid entered as an element into the composition of atmospherical air; and Bergman, adopting the same opinion, gave it the name of aerial acid. Mr Bewdly called it mepbitic acid, because it could not be respired without occasioning death; and this name was also adopted by Morveau. Mr Keir called it calcareous acid; and at last Mr Lavoisier, after discovering its composition, gave it the name of carbonic acid gas.

For the investigation of the properties of carbonic acid we are chiefly indebted to the labours of Caven-

^{*} Keit's Macquer, art. Air

ish , Priestley t, Bergman I, and Lavoisier §. Va- Chap. IL mable dissertations on it were also published by Macbride ||, Lane ¶, Jacquin **, Landriani ++, Fontana, and many other distinguished chemists.

1. Carbonic acid, being a compound of carbon and Preparabaygen, may be formed by burning charcoal; but as it exists in great abundance ready formed, there is no ocession to have recourse to that expedient. All that is ecessary is to dilute with water a quantity of chalk, which is a compound of carbonic acid and lime, and to pour upon it sulphuric acid. An effervescence ensues, carbonic acid is evolved in the state of gas, and may be received in the usual manner.

2. Carbonic acid, when in the state of gas, is invi- Properties. able and elastic like common air. It extinguishes a andle, and is unfit for respiration. It has no smell.

Its specific gravity, according to the experiments of Kirwan, is 1.500, that of air being 1.000; or it is to mir as 3 to 2. At the temperature of 60°, barometer 30 inches, 100 cubic inches of this gas weigh 46.5 grains 11. From this considerable weight, compared to that of air, it happens that this gas may be poured from one vessel to another. When poured from a widemouthed vessel upon a burning candle, it extinguishes it like water.

Phil. Trans. 1766, lvi. 141.

[†] Prientley, i. 43.

[🖁] Opasza i. L.

Mem. Par 1776.

Experimental Entrys, 1764.

Priestley, 1. 5

Diet. Meyer, de Acido Pingui, et Blachiana de Acido me, 1769.

¹¹ Crell's Amali, 1785, ii. 139.

^{##} Kirwan on Phloguton, p. 29

Pol. 11.

It reddens the tincture of turnsol, but no other vegetable colour *.

Atmospheric air is supposed to contain about reas part of this gas †.

3. Carbonic acid is not altered by exposure to heat in close vessels, or by passing it through a red hat tube. But when electric sparks are made to pass through it for a long time, its bulk is increased, as Dr Priestley first observed; and in that case, as Monger discovered, an inflammable gas is always evolved. By 18 hours labour Mr Saussure junior produced an increase of the part of the whole; one-tenth of the carbonic acid had disappeared, and a quantity of carbonic oxide had been formed. The copper wires employed to transmit the sparks were oxidized at the expence of the acid, which was thus partially converted into carbonic oxide so.

Action of water.

4. Water absorbs it gradually when allowed to remain long in contact with it. The rapidity of the absorption, as Mr Lane first discovered, is much increased by agitation. At the temperature of 41° water absorbs its own bulk of this gas. The specific gravity of water saturated with it is 1.0015. This water, at the temperature of 35°, has little taste; but if it be left a

[·] Bergman, i. 9.

[†] At least near the surface of the earth. Lamanon, Mongez, and the other unfortunate philosophers who accompanied La Perouse in his last voyage, have rendered it not improbable, that at great heights the quantity of this gas is much smaller. They could detect none in the atmosphere at the summit of the Peak of Teneriffe.—See Lamanon's Memoir at the end of La Perouse's Voyage.

[§] Fbil. Mag. xiv. 250.

few hours in the temperature of 88°, it assumes an Chap. II. agreeable acidity, and a sparkling appearance *.

Ice absorbs no carbonic acid; and if water containing. it be frozen, the whole separates in the act of freezing †. This gas also separates from water at the boiling temperature 1.

By means of artificial pressure, the quantity of this gas absorbed and condensed by water is much increased. Some of the aerated alkaline water, prepared in Britain as a medicine, is said to contain about thrice its volume of carbonic acid gas \.

When this liquid carbonic acid is left exposed to the air in an open vessel, the acid soon makes its escape in the form of gas, and leaves the water in a state of pu-Bergman left a bottle filled with carbonic acid rity. gas uncorked, and found that in a few days it contained nothing but common air.

5. Carbonic acid is not acted upon by oxygen, nor Action of is it altered by any of the simple combustibles, incombustibles, or metals. Charcoal indeed absorbs it, but it gives it out again unchanged. From the experiments of Rouppe and Van Noorden, it appears that this absorption is exceedingly rapid, provided the charcoal be sufficiently freed from air, by allowing it to cool from a red heat in a vacuum. They found that charcoal is capable of absorbing rather more than fourteen times its bulk of carbonic acid gas ¶. These experiments have been confirmed by others made in a different way by

simple box

Bergman, i. 9. † Priestley, i. 120. ‡ Ibid.

⁶ A very convenient apparatus for this purpose has been contrived by Mr Gilbert Austin, and described in the Isisb Transactions, viii. 131.

Ann. de Chim. XXXII. 11.

Count Morozzo". At a red heat charcosl converts into carbonic oxide gas.

Phosphorus is insoluble in carbonic acid gas †; but, phosphorus is capable of decomposing this acid by compound affinity, when assisted by a sufficient heat, as has been demonstrated by Tennant and Pearson. It on also and zinc, and several other metals, are capable of producing the same effect, as has been discovered by Priest-ley and Cruikshanks. In the first case the phosphorus is partly acidified, partly combined with lime, and charcoal is deposited: in the second, the metals are oxidized, and the gaseous oxide of carbon evolved.

When carbonic acid is mixed with sulphureted, phosphureted, or carbureted hydrogen gas, it renders them less combustible, but produces no farther sensible change.

Carbonates.

o. This acid is capable of combining with alkalies, and with several of the earths and metallic oxides, and forms with them salts, distinguished by the name of carbonates. All the earthy carbonates are insoluble in water. Hence the reason that carbonic acid renders lime water, barytes water, and strontian water turbid. The earth held in solution in these waters combines tapidly with the carbonic acid, which is of course absorbed, if in the state of gas, while the carbonate formed precipitates slowly in the form of a white powder. This occasions the milkiness which immediately ensues. As carbonic acid is the only gas not instantaneously absorbable by water, which renders lime was

^{*} Nicholson's Journal, iz. 262.

[†] Foureroy and Vauquelin, Ibid. zzi. 205.

ber, or barytic or strontian water, turbid, its presence Chap. Il. may be always ascertained by means of these liquids. As it is completely absorbed by them, it may be easily separated by them from other gases with which it may be mixed, and its quantity ascertained by the diminution of volume which the residuum of gas has undergone.

7. From the experiments of Saussure, it appears that carbonic acid scarcely combines with alumina *.

Water containing a little soda, and supersaturated with carbonic acid, has been employed with much advantage under the name of aerated alkaline water, as a palliative in cases of urinary calculus.

8. The opinious of chemists concerning the composi- Composition of carbonic acid have undergone as many revolutions as its name. Dr Priestley and Bergman seem at first to have considered it as an element; and several celebrated chemists maintained that it was the acidifying principle. Afterwards it was discovered to be a compound, and that oxygen gas was one of its compo-nent parts. Upon this discovery the prevalent opinion of chemists was, that it consisted of oxygen and phlogiston; and when hydrogen and phlogiston came (according to Mr Kirwan's theory) to signify the same thing, it was of course maintained that carbonic acid was composed of oxygen and hydrogen: and though Mr Lavoisier demonstrated, that it was formed by the combination of carbon and oxygen, this did not prevent the old theory from being maintained; because carbon was itself considered as a compound, into which a very

great quantity of hydrogen entered. But after Mr Lavoisier had demonstrated, that the weight of the carbonic acid produced was precisely equal to the charcoal and oxygen employed; after Mr Cavendish had discovered that oxygen and hydrogen when combined did not form carbonic acid, but water-it was no longer possible to hesitate that this acid was composed of carben and oxygen. Accordingly all farther dispute about it seems now at an end. At any rate, as we have already examined the objections that have been made to this conclusion, it would be improper to enter upon From the experiments of N:r Smithson them here. Tennant, compared with those of Lavoisier, and Messas Allen and Pepys, we may consider carbonic acid as composed of about

> 28 carbon 72 oxygen

100

Pecomposi-

If any thing was still wanting to put this conclusion beyond the reach of doubt, it was to decompound carbonic acid, and thus to exhibit its component parts by analysis as well as synthesis. This has been actually done by the ingenious Mr Tennant. Into a tube of glass he introduced a bit of phosphorus and some carbonate of lime. He then scaled the tube hermetically, and ap-Phosphate of lime was formed, and a plied heat. quantity of charcoal deposited. Now phosphate of lime is composed of phospheric acid and lime; and phosphoric acid is composed of phosphorus and oxygen. The substances introduced into the tube were phosphorus, lime, and carbenic acid; and the substances found in it were phosphorus, lime, oxygen, and charcoal. e carbonic acid, therefore, must have been decomed, and it must have consisted of oxygen and charl. This experiment was repeated by Dr Pearson,
o ascertained that the weight of the oxygen and
arcoal were together equal to that of the carbonic acid
ich had been introduced: and in order to show that
was the carbonic acid which had been decomposed,
introduced pure lime and phosphorus; and instead
obtaining phosphate of lime and carbon, he got nong but phosphuret of lime. These experiments*
re also confirmed by Messrs Fourcroy, Vauquelin,
lvestre, and Broigniart †.

SECT. VI.

OF FLUORIC ACID.

me mineral called fluor or fusible spar, and in this Mistory.

untry Derbysbire spar, was not properly distinguishfrom other spars till Margraff published a dissertan on it in the Berlin Transactions for 1768. He
st proved that it contained no sulphuric acid as had
en formerly supposed; he then attempted to decom-

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¹ Ann. de Chim. xiii. 382.

Count Mussiu-Puschkin having boiled a solution of carbonate of ash on purified phosphorus, obtained charcoal. This he considered in instance of the decomposition of carbonic acid, and as a confirmation the experiments related in the text. See Ann. de Chim. xxv. 105.

pose it by mixing together equal quantities of this mineral and sulphuric acid, and distilling them. By this method he obtained a white sublimate, which he supposed to be the fluor itself volatilized by the acid. He observed with astonishment that the glass retort was corroded, and even pierced with holes. Nothing more was known concerning fluor till Scheele published his experiments three years after; by which he proved that it is composed chiefly of lime and a particular acid, which has been called fluoric acid.

It is always obtained from fluor spar, in which mineral it is found in abundance. For the investigation of the properties of this acid, we are indebted chiefly to Scheele and Priestley.

Prepara-

1. It may be obtained by putting a quantity of the spar in powder into a retort, pouring over it an equal quantity of sulphuric acid, and then applying a very gentle heat. A gas issues from the beak of the retort, which may be received in the usual manner in glass jars standing over mercury. This gas is fluoric acid.

The acid may be obtained dissolved in water by luting to the retort a receiver containing water. The distillation is to be conducted with a very moderate heat, not only to allow the gas to condense, but also to prevent the fluor itself from subliming. After the process, provided a glass retort has been employed, a crust of white earth is found in the receiver, which has all the properties of silica.

^{*} In order to obtain this acid pure, platinum or silver vessels ought to be employed; for the acid dissolves a portion of glass and even of leaded vessels.

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Scheele supposed that the silica produced was formed of fluoric acid and water; and Bergman adopted the same opinion. But Wiegleb and Bucholz showed that the quantity of silica was exactly equal to what the retort lost in weight, and Meyer completed the proof that it was derived from the glass, by the following experiment. He put into each of three equal cylindrical tin vessels a mixture of three ounces of sulphuric acid and one ounce of fluor, which had been pulverized in a mortar of metal. Into the first he put one ounce of pounded glass; into the second, the same quantity of quartz in powder; and into the third, nothing. Above each of the vessels he hung a sponge moistened with water; and having covered them, he exposed them to a moderate heat. The sponge in the first cylinder was covered with the crust in half an hour, the sponge in the second, in two hours; but no crust was formed in the third, though it was exposed several days. In consequence of this decisive experiment Bergman gave up his opinion, and wrote an account of Meyer's experiment to Morveau, who was employed in translating his works, to enable him to correct the mistake in his notes.

Soon after the discovery of this acid, difficulties and doubts concerning its existence as a peculiar acid were started by some French chemists, disguised under the name of Boulanger, and afterwards by Mr Achard and Mr Monnet. To remove these objections Mr Scheele instituted and published a new set of experiments; which not only completely established the peculiar nature of the fluoric acid, but once more displayed the agrivalled abilities of the illustrious discoverer. It

[·] Scheele, ii. 38, and Crell's Annals, 1786, i. 3

would be needless to enumerate these objections, as they originated entirely from want of precision, and did not produce a single convert.

Properties.

- 2. Fluoric acid gas is invisible and elastic like air; it does not maintain combustion, nor can animals breathe it without death. It has a pungent smell, not unlike that of muriatic acid. It is heavier than common air. It corrodes the skin almost instantly.
- 3. Neither caloric nor light produce any alteration on it.

Action of water.

4. When water is admitted in contact with this gas, it absorbs it rapidly; and if the gas has been obtained by means of glass vessels, it deposits at the same time a quantity of silica.

Water absorbs a considerable proportion of this gas, but the precise quantity has not been determined. The compound is usually termed fluoric acid by chemists. It is specifically heavier than water, has an acid taste, reddens vegetable blues, and does not freeze till cooled down to 23°*. When heated, the acid gas is easily expelled, except the last portions of it, which adhere with great obstinacy.

Of simple bodies.

- 5. Neither oxygen gas nor any of the simple combustibles or incombustibles produce any change on fluoric acid, either in the gaseous or liquid state.
- tals; but liquid fluoric acid is capable of oxidizing iron, zinc, copper, and arsenic. It does not act upon gold, platinum, silver, mercury, lead, tin, antimony, cobalt.

Fluates

7. It combines with alkalies, earths, and metallic ox-

^{*} Priestley, ii. 361.

des, and forms with them salts which are denominated

The most singular property of fluoric acid is the fathity with which it corrodes glass and siliceous bodies, especially when bot, and the ease with which it holds illica in solution even when in the state of gas. This Unity for silica is so great, that the thickest glass vesels can only withstand its action for a short time; and the greatest precautions are scarcely sufficient to obtain B entirely free from siliceous earth.

S. It produces no change, as far as is known, upon my of the acids already described.

9. As fluoric acid produces an insoluble compound with lime, it may be employed with great advantage, Pelletter has observed, to detect the presence of that earth when held in solution. A drop or two of the heid causes a milky cloud or precipitate to appear, if my lime be present. The property which this acid has Used for corroding glass, has induced several ingenious men glass. attempt, by means of it, to engrave, or rather etch spon glass. The glass is covered completely with wax; and then that part where the letters or figures are to appear is laid bare by removing the wax. The whole is then exposed for some time to the hot vapours of fluoric acid. This simple process is employed with advantage in writing labels on glass vessels, and in graduating thermometers, and other similar instruments. The discovery is by no means new: It has been shown by Beckman and Accum, that this acid was employed for bat purpose by Henry Swanhard, an artist of Nuremberg, as early as 1670. He seems to have kept his

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art for some time secret; but the receipt was made public by Pauli in 1725*.

Till lately we were entirely ignorant of the constituents of this acid. Dr Henry tried in vain to decompose it by means of electricity; but Mr Davy's late attempts have been more successful. Potassium has the property of decomposing it, and the decomposition is attended with combustion †. A proof that oxygen is one of its constituents. The base of the acid is analogous to sulphur ‡. It combines with the potash formed, or with the potassium if there be an excess of that substance. It is combustible, and produces by its combustion fluoric acid. Hence it is obvious that this acid is truly an acid product. No name has hitherto been given to its peculiar basis.

^{*} See Beckman's History of Inventions, and Accum, Nicholson's Jannel. iv. 1.—The following advertisement, published by these gentlement from a German publication of 1725, will give an idea of the method enployed by that artist. " Take spiritus nițri per distillationem, put it into a retort, and apply a strong heat. When it has passed over into the receiver, throw into it some powdered green Bebenius emerald (which, when heated, skines in the dark), otherwise called hophorus. This being done, place the receiver, containing the mixture, on a heated sadbath for about four-and-twenty hours, and it will be fit for the purpose. To use this corrosive acid, take a pane of glass of any kind, clean it well, and free it from grease by washing with a ley; and when dry, trace est upon it, with sulphur and varnish, whatever you choose. Put a border of bees wax round it about one inch high, and then pour the correive acid, prepared as before directed, carefully over the whole surface of the glass, and let it stand undisturbed for some time; the longer the better. The glass will become corroded; and all you had traced before will now appear as if raised, or elevated, above the surface of the glass, in a very distinct and pleasing manner." The hesphorus here mentioned was evidently fluor spar.

[†] Pbil. Meg. xxiii. 89.

[†] Nicholson's Journal, 221. 238.

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SECT. VII.

OF BORACIC ACID.

Arabian chemist of the 10th century. It is a name wen to a species of white salt much used by various tists. Its use in soldering metals appears to have sen known to Agricola.

Borax is found mixed with other substances in Thi-It seems to exist in some lands adjacent to lakes, from which it is extracted by water, and deposited in bose lakes; whence in summer, when the water is Mallow, it is extracted and carried off in large lumps. smetimes the water in these lakes is admitted into reervoirs; at the bottom of which, when the water is shaled by the summer's heat, this salt is found. Hence is carried to the East Indies, where it is in some measure purified and crystallized : in this state it comes Europe, and is called tineal. In other parts of Thiet, it seems, by accounts received from China, they it out of the ground at the depth of about two yards, where they find it in smaller crystalline masses, called y the Chinese mi poun, bour poun, and pin pouin; and e earth or ore is called pounza *.

Though borax has been in common use for nearly tree centuries, it was only in 1702 that Homberg, by

[·] Kirwan's Miner. ii. 3"

vered the boracic acid. He called it sarcotic or sedative salt, from a notion of his that it possessed the properties indicated by these names. In his opinion, it was merely a product of the vitriol which he had used; but Lemery the Younger soon after discovered, that it could likewise be obtained from borax by means of the nitric and muriatic acids. Geoffroy afterwards discovered that borax contained soda; and at last Baron proved, by a number of experiments, that borax is composed of boracic acid and soda; that it may be reproduced by combining these two substances; and that therefore the boracic acid is not formed during the decomposition of borax, as former chemists had imagined, but is a peculiar substance which pre-existed in that salt.

This conclusion has been called in question by Mr Cadet *; who affirmed, that it was composed of sode, the vitrifiable earth of copper, another unknown metal, and muriatic acid. But this assertion has never been confirmed by a single proof. Mr Cadet has only proved, that boracic acid sometimes contains copper; and Bauné's experiments are sufficient to convince us, that this metal is merely accidentally present, and that it is probably derived from the vessels employed in crystallizing borax; that boracic acid generally contains a little of the acid employed to separate it from the soda with which it is combined in borax; and that crude borax contains a quantity of earth imperfectly saturated with boracic acid. All which may be very true; but they are altogether insufficient to prove that boracic acid is

⁴ Jour. de Phys. 1782.

not a peculiar substance, since it displays properties different from every other body.

Messrs Exschaquet and Struve * have endeavoured, on the other hand, to prove, that the phosphoric and boracic acids are the same. But their experiments merely show, that these acids resemble one another in several particulars; and though they add considerably to our knowledge of the properties of the phosphoric acid, they are quite inadequate to establish the principle which these chemists had in view; since it is not sufficient to prove the identity of the two acids, to show us a resemblance in a few particulars, while they differ in many others. Boracic acid must therefore be considered as a distinct substance.

- 1. The easiest method of procuring boracic acid is Preparathe following one: Dissolve borax in hot water, and filter the solution; then add sulphuric acid, by little and little, till the liquid has a sensibly acid taste. Lay it aside to cool, and a great number of small shining laminated crystals will form. These are the boracic acid. They are to be washed with cold water, and drained upon brown paper.
- 2. Boracic acid, thus procured, is in the form of thin Properties. hexagonal scales, of a silvery whiteness, having some resemblance to spermaceti, and the same kind of greasy It has a sourish taste at first +, then makes a bitterish cooling impression, and at last leaves an agreeable sweetness. It has no smell; but when sulphuric acid

[#] Jour. de Phys. xxviii. 1 16.

[†] Owing most probably to the remains of the acid employed in procoring it; for it loses that taste when heated to redness.

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is poured on it, a transient odour of musk is produced.

It reddens vegetable blues. Its specific gravity is 1.479 + while in the form of scales; after it has been fused it is 1.803 ‡.

3. It is not altered by light. It is perfectly fixed in the fire. At a red heat it melts, and is converted into a hard transparent glass; which becomes somewhat opaque when exposed to the air, but does not attract moisture.

Action of water,

4. It is much less soluble in water than any of the acids hitherto described. Boiling water scarcely dissolves 0.02 of boracic acid, and cold water a still smaller quantity. When this solution is distilled in close vessels, part of the acid evaporates along with the water, and crystallizes in the receiver. Water, therefore, renders it in some measure volatile, though it is perfectly fixed when in a state of dryness.

Of simple bodies.

- 5. Neither oxygen gas, the simple combustibles, the simple incombustibles, nor the metals, produce any change upon boracic acid, as far as is at present known.
 - 6. It is soluble in alcohol; and alcohol containing it burns with a green flame. Paper dipped into a solution of boracic acid burns with a green flame.

Though mixed with fine powder of charcoal, it is nevertheless capable of vitrification; and with soot it melts into a black bitumen-like mass, which is, however, soluble in water, and cannot be easily calcined to ashes, but sublimes in part §.

^{*} Reuss, de Sala Sedat. 1778.

[†] Kirwan's Min. ii, 4.

¹ Ha-senfratz, Ann. de Clim. xxviii. 11.

[&]amp; Keit's Dictionary.

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the assistance of a distilling heat, it dissolves in initially in mineral oils; and with these it yields solid products, which give a green colour to wine.

boracic acid is rubbed with phosphorus, it prevent its inflammation; but an earthy yelster is left behind *.

hardly capable of oxidizing or dissolving any of als except iron and zine, and perhaps copper. ic acid combines with alkalies, alkaline earths, mina, and most of the metallic oxides, and forms

mas which are called borates.

component parts of this acid were till lately un- Composi-Fabroni announced that he considers it as a mon of muriatic acid, and that it may be preom that acid; but he has not yet published the ents upon which these opinions are founded +. perious set of experiments have been made on it

By digesting oxymuriatic acid on it for a time, he succeeded in decomposing it, and from it a substance exactly resembling charthe properties, and a volatile acid resembling the in the greater number of its properties, but from it in not precipitating lead from its solu-Mr Davy has succeeded in decomposing this means of potassium, and has shown that it is and of oxygen, and a combustible basis of a black

eII.

Dictionary. † Foureroy, ii. 128. dim. xxxv. 202.—Crell considers this volatile acid as very the rebarn.

colour, which has a greater resemblance to charcoal than to any other substance. It becomes white, and is doubtless converted into boracic acid by exposure to the air.

• Nichelson's Jeer. xxii. 238.

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CLASS II.

ACID SUPPORTERS.

acid supporters are distinguished by the follow- properties:

They cannot be produced by combustion. Hence pase is either a simple incombustible or a metallic

They are capable of supporting combustion. Hence eadily acidify the combustible bases, and convert of the metals into oxides.

They are decomposed by exposure to a high temre; their oxygen in that case making its escape state of a gas.

e only acids which possess these properties in pernare those which have the simple incombustibles resenic for their bases; but from analogy I refer hole of the metallic acids to this head. None of can be produced by combustion; and as they n a maximum of oxygen, they are of course innatible: They cannot, therefore, be referred to any other two classes.

The following TABLE exhibits a view of all the scill supporters, their composition, and the proportion of their constituents, as far as they have been ascertained.

Number and composition.

Acids.	Bases.	Propertion of Oxygen to 100 Base.	
Nitric Nitrous	Azote	236	
Oxymuriatic Hyperoxymuriatic	Muriatic seid	29 19 4	
Arsenic	Arsenic	53	
Tungstic	Tungsten	25	
Molybdic	Molybdenum	59	
Chromic	Chromium	200	
Columbic	Columbium		

Some of these acids are of great importance in chemistry. This is chiefly owing to their activity: for which they are indebted to the state of the oxygen which they centain. The theory of Lavoisier applies with precision to this class of acids as well as to the first.

The second secon

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SECT. I.

OF WITRIC ACID.

acro seems to have been first obtained in a History. atate by Raymond Lully, who was born at Ma-1235. He procured it by distilling a mixture and clay. Basil Valentine, who lived in the stury, describes the process minutely, and calls water of nitre. It was afterwards denominafortis and spirit of nitre. The name nitric acid t given it in 1787 by the French chemists. is generally obtained in large manufactories by Preparag a mixture of nitre and clay; but the acid proby this process is weak and impure. Chemists y prepare it by distilling three parts of nitre of sulphuric acid in a glass retort. This mea first used by Glauber. The neck of the renet be luted into a receiver, from which there glass tube into a bottle with two mouths, cona little water, and furnished with a tube of safe-From the other mouth of this bottle there passes

whe of sefety is a tube open at its upper end, and having its plunged in water. The water prevents any communication the external air and the inside of the apparatus. If a vacuum to be formed within the vessels, the external air rushes down the tube and prevents any injury to the vessels. On the other air is generated in the vessels, it forces the water up the tube, but of which becomes thus the measure of the elasticity of the air mels. By this contrivance the apparatus is in no danger of below, which otherwise might happen.

a tube into a pneumatic apparatus to collect the gas which is evolved during the process. The apparatus is represented in fig. 12. The retort is to be heated gradually almost to redness. The nitric acid comes over and is condensed in the receiver, while the common air of the vessels, and a quantity of oxygen gas which is evolved, especially towards the end of the process, passes into the pneumatic apparatus, and the water in the bottles is impregnated with some acid which is not condensed in the receiver.

The acid, thus obtained, is of a yellow colour, and almost always contains muriatic and sulphuric acid, from which it is difficult to free it completely. Perhaps the best way is to purify the nitre beforehand by repeated crystallizations, and by throwing down the muriatic acid that may adhere after all, by means of nitrate of silver. From the experiments of Lassone and Cornette, it appears that if nitric acid contaminated with muriatic be distilled with precaution, the whole of the muriatic acid comes over with the first portions, and the last portions are quite free from it *. The common method is to mix impure nitric acid with nitrate of silver, to separate the precipitate, and re-distil. This method succeeds only when the acid is strong; if it be weak, a portion of muriatic acid still adheres to it. The sulphuric acid may be separated by rectifying the acid, by distilling it slowly, and withholding the last portions, or by distilling it off litharge or nitrate of barytes.

After these foreign bodies are separated, the acid still retains a quantity of nitrous gas, to which it owes its

^{*} Mem. Per. 1781, p. 645.

four and she red fumes which it exhales. This gas Chap IL by be expelled by the application of heat. Pure sitric remains behind, transparent and colourless, like ster.

When newly prepared in this manner, it is a li- Properties. id as transparent and colourless as water; but the Buity between its component parts is so weak, that the Mon of light is sufficient to drive off a part of its oxyin the form of gas; and thus, by converting it intly into nitrous gas, to make it assume a yellow comr. Its taste is exceedingly acid and peculiar. It is by corrosive, and tinges the skin of a yellow colour, hich does not disappear till the epidermis comes off. is constantly emitting white fumes, which have an aid and disagreeable odour.

. It has a strong affinity for water, and has never yet Action en obtained except mixed with that liquid. eentrated, it attracts moisture from the atmosphere, not so powerfully as sulphuric acid. It also probes heat when mixed with water, owing evidently to concentration of the water.

The specific gravity of the strongest nitric acid that be procured is, according to Rouelle, 1'583; but the temperature of 60°, Mr Kirwan could not proit stronger than 1-5543.

But as this liquid acid is a compound of two ingre- Of the ts, namely, pure nitric soid and water, it becomes nitric soid. object of the greatest consequence to ascertain the aportion of each of these parts. This problem has My occupied the attention of Mr Kirwan, who has deavoured to solve it in the following manner.

He dried a quantity of crystallized carbonate of soda ared heat and dissolved it in water, in such a pro-

portion that 367 grains of the solution contained 50.05 of alkali. He saturated 367 grains of this solution with 147 grains of nitric acid, the specific gravity of which was 1.2754, and which he ascertained to contain 45.7 per cent. of acid, of the specific gravity 1:5543. chosen by him as a standard. The carbonic acid driven off amounted to 14 grains. On adding 939 grains of water, the specific gravity of the solution, at the temperature of 58.5°, was 1.0401. By comparing this with a solution of nitrate of soda, of the same density, precisely in the manner described formerly under sulphuric acid, he found, that the salt contained in it amounted to 10:001 of the whole. There was an excess of acid of about two grains. The weight of the whole was 1439 grains: The quantity of salt coasequently was 1439 = \$5.142 grains. The quantity of alkali was 50.05 - 14 = 36.05. The quantity of standard acid employed was 67.18; the whole of which amounted to 103.23 grains: but as only 85.142 grains entered into the composition of the salt, the remaining 18.088 must have been pure water mixed with the nitric acid. But if 67.18 of standard acid contain 18.088 of water, 100 parts of the same acid must contain 26.92 *.

One hundred parts of standard nitric acid, therefore, are composed of about 73.62 parts of pure nitric acid, and 26.38 of water. But as Mr Kirwan has not proved that nitrate of soda contains no water, perhaps the

portion of water may be greater. He has rendered it bable, however, that nitrate of soda contains very de water.

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Mr Kirwan's real acid, then, is nitric acid, of that tree of strength which enters into the composition of trate of soda. The proportion of this real acid, conmed in nitric acid of different specific gravities, has an given by Mr Kirwan in the following TABLE:

co Parts	Real 100 Parts		Real	
Sp. Gravity.	Acid.	Sp Gravity.	Acid.	
215540	Co. A		7.5	
1.5543	31-4	19975	50*74	
1.5295	69 86	113925	50100	
1.2183	69.12	1.3875	49.21	
1.2010	68-39	1.3825	48151	
1.4957	67-65	1.3715	47.80	
1.4844	60.85	1.3721	47'06	
1:4731	66*18	1.3671	46.33	
1.4710	65*45	1.3621	45159	
1.4707	64.71	1.3571	44.86-	
1'4695	63 98-	1.3521	44-12	
1.4683	63:24	1.3468	43*38	
114071	02:51	1.3417	42.65	
1:4640	61.77	1.3364	41-91	
1:4611	61.03	1.3315	4P18	
1.4582	60.30	1.3264	40*44	
1-4553	59.56	1.3212	39.71	
1.4524	58.83	1.3160	38'97	
1:4471	58.09	1'3108	38-34	
1.4422	57'36	1.3056	37.50	
1-4373	56.62	1.3004	36.77	
1.4324	55.89	1.2911	36.03	
1.4275	55.15	1.2812	35.30+	
1.4222	54.12+	1.2795	34.20	
1.4171	53.68	1.2779	33.82	
1.4120	52.94			
1-4069	52.21	1.2687	31.09	
1.4018	51. 7	1:2586	32135	
2 4013	J 2 1	1.500	1'62	

100 Parts Sp. Gravity.	Real Acid.	100 Carts Sp. Gravity.	Real Acid.
1.2464	30.88	1.1779	22.06
1.2419	30.12	1.1201	21.32
1.2374	29.41	1.1039	20.20
1.2291	29.68	11581	19.85
1.2209	27.94	1.1524	19.12
1.2180	27.21+	1.1421	18.48
1.2152	26.47	1.1319	17.65+
1.2033	25.74+	1.1284	16.91
1.2015	.25.00	1.1241	16.17
1.1963	24.26	1.1165	15.44
1.1911	23.53	1.1111	14.70
1.1845	24.79	1.1040	13.27

Mr Davy considers as pure acid the permanently elastic vapour or gas formed by saturating nitrous gas with oxygen gas. This gas is of a pale yellow colour, and a specific gravity 2.44 times that of air. It is not pure acid, containing undoubtedly a portion of nitrous gas. The following TABLE exhibits the proportion of this acid contained in nitric acid of different densities, according to the experiments of that ingenious chemist.

Nitric acid, of Sp. Gr.	True acid.	Water.
1.5040	91.22	8.45
1.4475	80.38	19.61
1.4285	71.65	28.35
1.3906	62.96	37.0+
1.3551	56.88	43.12
1.3186	52.03	47.07
1.3042	49.04	50.96
1.5831	40.03	53.97
1.5000	45.27	54.13

[#] Duvy's Researches, p. 41.

4. When pitric acid is exposed to the action of heat, boils at the temperature of 24800, and evaporates cometely without alteration; but when made to pass brough a red hot porcelain tube, it is decomposed, and enverted into oxygen and azotic gas †. When cooled own to -66°, it begins to congeal; and when agitait is converted into a mass of the consistence of but-1. But an account of the freezing points of this aid, as ascertained by Cavendish, has been already Iven in a preceding part of this Work f. Lassone and Cornette have ascertained, that when weak nitric acid boiled or distilled, the weakest portion comes first wer into the receiver; but when the acid is conceninted, the strongest portion comes first over | .

5. Oxygen gas has no action whatever on nitric acid; Action of all the simple combustibles decompose it, unless stances. we are to except the diamond. When poured upon alphur or phosphorus \ at a high temperature, it sets em on fire; but at a moderate temperature it conserts them slowly into acids, while nitrous gas is exled. It inflames charcoal also at a high temperature. ad even at the common temperature, provided the charcal be perfectly dry and minutely divided **. Hydroen gas produces no change on it at the temperature of atmosphere; but when passed along with it through

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Bergman, it. 141. † Fourcroy, ii. 82. Fourcioy and Vauquelin, Ann. de Chim. xxxx. 281.

Wel. 1 p. 523. 1 vem. Per. 1781, p. 645.

According to Brugnatelli, when a bit of phosphorus is wrapt up in per, dept in nitric soid, and struck smartle with a hammer upon an wil, a detonation takes place Jour. de Chim, iv. 113.- With me, howas Proust. the experiment did not succeed.

a red hot porcelain tube, it detonates with great violence; water is formed, and azotic gas evolved *.

When this acid is poured upon oils, it sets them on This is occasioned by a decomposition both of the acid and oil. The oxygen of the acid combines with the carbon and with the hydrogen of the oils, and at the same time lets out a quantity of caloric. Hence we see that the oxygen which enters into the composition of the nitric acid still contains a great deal of caloric; a fact which is confirmed by a great number of other The combustion of oils by this acid was phenomena. first taken notice of by Borrichius and Slare +; but it is probable that Homberg communicated it to Slare. In order to set fire to the fixed oils, it must be mixed with some sulphuric acid; the reason of which seems to be, that these oils contain water, which must be previously removed. The sulphuric acid combines with this water, and allows the nitric acid, or rather the oil and nitric acid together, to act. The drying oils do not require any sulphuric acid: they have been boiled, and consequently deprived of all moisture.

- o. Azote has no action on nitric acid; but muriatic acid decomposes it by combining with a portion of its oxygen, nitrous gas and oxymuriatic gas being evolved. A mixture of nitric and muriatic acids was formerly called aqua regia; it is now known by the name of nitro-muriatic acid.
- 7. It is capable of oxidizing all the metals except gold, platinum ‡, and titanium. It appears, from the

^{*} Fourcroy, ii. 82. † Phil. Trans. Abr. ii. 653, and iii. 663. † Nitre, however, acts upon platinum, as Mr Tennant has proved. Phil. Trans. 1797.—Morveau had made the same observation in the Elimens de Chimie de l' Academie de Dijon.

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experiments of Scheffer, Bergman, Sage, and Tillet. that nitric acid is capable of dissolving (and consequently of oxidizing) a very minute quantity even of old.

It even sets fire to zinc, bismuth, and tin, if it be poured on them in fusion, and to filings of iron if they he perfectly dry *.

8. Nitric acid combines with alkalies, earths, and the exides of metals, and forms compounds which are call-Ma mitrates.

9. Nitric acid absorbs nitrous gas with great avidity. assumes a yellow or brown colour, and the property of bmitting dense yellow fumes. In this state it was formerly known by the name of phlogisticated nitric acid.

The liquid at present distinguished in most chemical books by the name of nitrous acid, is nothing else than mis combination. Its nature was first investigated by Dr Priestley, who demonstrated, by very decisive experiments, that it is a compound of nitric acid and nifrous gas. This opinion was embraced, or rather it was wirst fully developed, by Morveau +. But the theory A comof Lavoisier, which supposed the difference between mitric and colourless and yellow nitric acid, to depend merely on and nitric the first containing a greater proportion of oxygen than the second, for some time drew the attention of chemists from the real nature of the combination. found published a dissertation in 1796, to demonstrate the truth of the theory of Priestley and Morveau; and the same thing has been done still more lately by Messrs Thomson and Davy t.

Proust, Dijon Academicians, and Cornette.

Mary: Merbod Chin. i. 18,

Davy's Researabes, p. 300

Nitric acid being capable of absorbing very proportions of nitrous gas, it is evident that the be a great variety of nitrous acids, to use the phrase, differing from each other in the propositrous gas which they contain; unless we confine the term to the compound formed by a nitric acid completely with nitrous gas.

When nitrous gas is placed in contact w. acid, the acid absorbs it slowly, and acquire pale yellow colour, then a bright yellow: Wh siderable portion more of nitrous gas is absor acid becomes dark orange, then olive, which in intensity with the gas absorbed; then it be a bright green; and lastly, when fully saturat comes blue green. Its volume and its volat increase with the quantity of gas absorbed; a fully saturated, it assumes the form of a dense of an exceedingly suffocating odour, and difficu densible by water. In this state of saturation tinguished by Dr Priestley by the name of sit vapour. It is of a dark red colour, and passes water partly without being absorbed. The qu nitrous gas absorbed by nitric acid is very gre Priestley found, that a quantity of acid, equal is four pennyweights of water, absorbed 130 ou eures of gas without being saturated *. The co parts of nitrous acid, of different colours and may be seen in the following TABLE, drawn u Davy, from experiments made by him on purp much precision +.

^{*} Priestley, i. 383.

100 Parts.	Sp. Grav	Component Parts.		
		Nitric acid.	Water.	Nitrous Gas.
Solid nitric acid	1.204	91.55	8.45	
Yellow nitrous	1.502	90*5	8*3	2
Bright yellow .	1.500	88.94	8.10	2193
Dark orange	1.480	86'84	7.6	5.56
Light olive	1.479	86100	7:55	6.45
Dark olive	1.478	85.4	7.5	7:1
Bright green	1.476	84-8	7-44	7.76
Bue green	1.475	84.6	7.4	8.00

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The colour of nitric acid depends, in some measure, also on the proportion of water which it contains. When to yellow nitric acid concentrated, a fourth part by weight of water is added, the colour is changed to a fine green; and when equal parts of water are added. il becomes blue *. Dr Priestley observed, that water impregnated with this acid in the state of vapour became first blue, then green, and lastly yellow. A green nitric acid became orange-coloured while hot, and reuned a yellow tinge when cold. A blue acid became yellow on being heated in a tube hermetically sealed. An orange-coloured acid, by long keeping, became green, and afterwards of a deep blue; and when exposed to air, resumed its original colour. When yellow nitric acid is exposed to heat, the nitrous gas is expelled, and nitric acid remains behind. however, carries along with it a quantity of acid, especially if the acid be concentrated. But nitrous acid vapour is not altered in the least by exposure to heat +.

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Nitrous acid vapour is absorbed by sulphuric aci but seemingly without producing any change; for who water is poured into the mixture, the heat produced expels it in the usual form of red fumes. The onlessingular circumstance attending this impregnation is, the it disposes the sulphuric acid to crystallize. This factivate observed by Dr Priestley in 17771, was afterward confirmed by Mr Cornette.

It is absorbed also rapidly by nitric acid, which as sumes the different colours which distinguish nitrou acid, according to the proportion which it imbibes.

10. Nitric acid is one of the most important instruments of analysis which the chemist possesses; nor is it of inferior consequence when considered in a political or commercial view, as it forms one of the most essential ingredients of guppowder. Its nature and composition accordingly have long occupied the attention of

Composition.

* Priestley, iii. 144 † Ibid. p. 156.

Pernhardt, however, relates, in 1765, that once, when he was disconsisted a maximum of ten pounds of nitre with an equal quantity of calcing vitriol, which he had put into a retort, to which he fitted an adopt between the actort and the receiver which contained a quantity of we ter—he observed a considerable quantity of a white crystalline salt formed in the adopter, while the liquid acid passed as usual into the receive. This salt was very volatile, smoked strongly when it was exposed to the air, and exhaled a red vapour'; it burnt to a black coal wood, feather or linen, as supplied acid does; and where a piece of it fell, it evapouted in form of a blood-red vapour, till the whole of it disappeared. He are conce of these crystals dissolved in water with spurting and hissing like that of a red hot iron dipped in water, and formed a green nitro acid. Some of this salt in a green bottle, which was not well stopped, entirely vanished. The analystals were evidently the same with I Priestley's. See Keir's Dictionary.

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ps by which its component parts were discovered.

As nitre is often produced upon the surface of the rth, and never except in places which have a commutation with atmospheric air, it was natural to suppose at air, or some part of the air, entered into the committee air, or some part of the air, entered into the committee and atmospherical air were both possessed of the reperty of giving a red colour to the blood, and that is was deprived of this property by combustion and espiration—concluded that nitre contained that part of the air which supported combustion, and was necessary in respiration.

Dr Hales, by applying heat to nitric acid, and what he called Walton mineral, obtained a quantity of air possessed of singular properties. When atmospherical hir was let into the jar which contained it, a reddish arbid finne appeared, a quantity of air was absorbed, and he remainder became transparent again. Dr Priestley discovered, that this air could only be obtained from nitric acid; and therefore called it hitrous air; He tound, that when this gas was mixed with oxygen gas, nitric acid was reproduced. Here, then, we find that exygen is a part of the nitric acid, and consequently that May ow's affirmation is verified.

Di Priest ey, however, explained this fact in a different manner. According to him, nitrous gas is composed of nitric said and phlogiston. When oxygen is added, it separates this phlogiston, and the acid of course is precipitated. This hypothesis was adopted by Mae-

[·] Veget. Statics, il. 284.

quer and Fontana; and these three philosophers ended voured to support it with their usual ingenuity. But there was one difficulty which they were unable to me mount. When the two gases are mixed in proper part portions, almost the whole assumes the form of nice acid; and the small residuum (1 th part), in all probability, or rather certainly, depends on some accident impurity in the two gases. What then becomes of the exygen and phlogiston? Dr Priestley supposed that they formed carbonic acid gas; but Mr Cavendish proved, that when proper precautions are taken, no such aid appears.

Dr Priestley had procured his nitrous gas by disch ving metals in nitric acid; during the solution of which a great deal of nitrous gas escapes. He supposed the nitrous gas contained phlogiston, because the metal wa exidized (and consequently, according to the then asceived theory, must have lost phlogiston) during its for Mr Lavoisier proved, that this supposite was ill founded, by the following celebrated experiment †. To 945 grains of nitric acid (specific gravity 1.316) he added 1104 grains of mercury. During the solution 273.234 cubic inches of nitrous gas were pro-He then distilled the salt (oxide of mercury) duced. which had been formed to dryness. As soon as it became red hot it emitted oxygen gas, and continued to do so till almost the whole of the mercury was revivel: The quantity of oxygen emitted was 287-749 cabie All that had happened, therefore, during the solution of the mercury, was the separation of the said

+ Men. 1776, p. 673

two parts; mitrous gas, which flew off, and oxygen; ch united with the metal.



Ar Lavoisier concluded, therefore, that the whole of sitrous gas was derived from the nitric acid; that ic weld is composed of oxygen and nitrous gas; and the proportions are nearly 64 parts by weight of our gas, and 36 of oxygen gas:

whedged he could not remove. The quantity of rgen obtained by decomposing nitric acid was often the greater than what was necessary to saturate the rous gas. Mr De Morveau attempted to account for spous without success. Nitrous gas itself was idently a compound: but the difficulty was to discontently a compound: but the difficulty was to discontently and periment made by decomposing nitre by means of mean, that it contained azote; and several of Driestley's experiments led to the same result. But hat was the other ingredient?

the Cavendish had observed, while he was making the composition of water, that some nicacid was formed during the combustion of oxygen and hydrogen gas, and that its quantity was increased yadding a little azote to the two gases before the exlosion. Hence he concluded, that the formation of he acid was owing to the accidental presence of azotic as. To verify this conjecture, he passed electrical parks through a quantity of common air inclosed in a lass tube: the air was diminished, and some nitric aid formed. He repeated the experiment, by mixing

^{*} Eneye. Metbed. Chim. Atide Nitrique

together oxygen and azotic gas; and found, that when they bore a certain proportion to each other they was totally convertible into nitric acid.

These experiments were immediately repeated by Messrs Van Marum and Van Treoctwyk, and with nearly the same result.

Dr Priestley had observed several years before the experiments were made, that atmosphesical air was a minished by the electric spark, and that during the aminution the infusion of turnsol became red; but he concluded merely that he had precipitated the acid of the air. Landriani, who thought, on the contrary, the carbonic acid gas was formed, enounced the alternian of lime-water by it as a proof of his opinion. It was to refute this notion that Mr Cavendish undertook his experiments.

It cannot be doubted, then, that nitrie acid is composed of azote and oxygen; consequently nitrons gas it also composed of the same ingredients. And as nitrous gas absorbs oxygen, even from common air, and forms with it nitrie acid, it is evident that nitrie acid contains more oxygen than nitrous gas. But it is exceedingly difficult to ascertain the exact proportions of the component parts of this acid. Lavoisier concluded, from his experiments on the decomposition of nitre by charcoals that nitric acid is composed of one part of azote and four parts of oxygen. But Davy has shown that this decomposition is more complicated than had been supposed; and that Lavoisier's experiments by at means warrant the conclusion which he drow from

Alem. de Sav. Etrang. 21. 226.

chap. II. Cavendish, on the other hand, concluded from the truth as nearly as possible. Nitric than is composed of 29.77 azote

70.23 oxygen

100.00

urly 1 part of azote to 21 of oxygen.

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OF NITROUS ACID.

note, which is a compound of nitric acid and h, is exposed to a red heat, it yields a considerable ity of oxygen gas almost in a state of purity. If socess be conducted with the proper precautions topped in time, the nitre still retains the properties sentral salt. But the acid which it contains is obly in a different state, since it has lost a consider-part of its oxygen. To this new state the term is acid is applied.

Davy's Researches, 1.44.

The experiment just recited was first made by Scheele, and mentioned in his dissertation on manganese, published in 1774 *. He first pointed out the difference between nitric and nitrous acids, but confounded nitrous acid with nitric acid impregnated with nitrous gas, His opinions were adopted by Bergman, and modified by Lavoisier to suit his own theory, Not with standing the experiments of Priestley, which explained the nature of fuming nitric acid in a satisfactory manner, the opinion of Lavoisier and Scheele prevailed, and the term nitrous was applied to nitric acid whenever it was co. loured with nitrous gas. Raymond drew the attention of chemists to the real constitution of furning nitric acidia 1796, and the same thing was done more lately by Messrs Davy and Thomson. It is now generally al. lowed, that it is merely a combination of nitric acid and nitrous gas; and as such, its properties have been detailed in the preceding Section.

But the nitrous acid formed by exposing nitre to heat is undoubtedly a different substance, since nothing like the new salt can be formed by uniting potash to fuming nitric acid. But all the attempts to separate the nitrous acid from the potash have failed. When an acid, however weak, is applied, fumes of nitrous vapour are immediately disengaged. For any thing we know to the contrary, nitrous acid can exist only combined with a base. The genus of salts which it forms are called mitrites. They are obtained by exposing the nitrates to a graduated heat for a certain time, and then stopping the process. None of them have been hitherto examined with attention except the nitrite of potash.

Scheele's Opuse. i. 59.

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SECT. III.

OF OXYMURIATIC ACID.

Enown, we may at present consider it as a simple substance. As it differs from all other acids in having the property of uniting with oxygen and forming a new set of acids which support combustion, it was thought preserable to separate it from the other acids, and to describe it along with azote, to which it bears a striking analogy. When united to oxygen, it forms the two acid supporters called oxymuriatic and hyperoxymuriatic; the properties of which we shall examine in this and the succeeding Section.

Oxymuriatic acid was discovered by Scheele in 1774, History. during his experiments on manganese. He gave it the name of dephlogisticated muriatic acid, from the supposition that it is muriatic acid deprived of phlogiston. The French chemists, after its composition had been ascertained, called it oxygenated muriatic acid; which unwieldy appellation Kirwan has happily contracted into exymuriatic.

The properties of this acid pointed out by Scheele were so peculiar, that it immediately attracted attention, and the most distinguished chemists hastened with emulation to enter upon a field which promised so rich a harvest of discoveries. Bergman, Pelletier, Berthollet, Hermbstadt, Morveau, Fourcroy, Scopoli, Westrumb, &c. successfully examined its properties, and ascertained its

action on other bodies; and more lately Mr Cheneviz has published a most interesting dissertation on this acid and its combinations.

Prepara-

1. It may be procured by the following process: Pat into a tubulated retort a mixture of three parts of common salt, and one part of the black oxide of manganese in powder. Place the retort in the sand bath of a furnace, plunge its beak into a small water trough, and lute a bent funnel into its mouth. When the mixture has acquired a moderate heat, pour into it at intervals through the bent funnel two parts of sulphuric acid, which ought to be somewhat diluted with water. An effervescence ensues, a yellow coloured gas issues from the retort, which may be received in large phials fitted with ground stoppers.

Properties.

- 2. Oxymuriatic acid gas is of a yellowish green colour. Its odour is intolerably acrid and suffocating. It cannot be breathed without proving fatal. The death of the ingenious and industrious Pelletier, whose chemical labours have been so useful to the world, was occasioned by his attempting to respire it. A consumption was the consequence of this attempt, which in a short time proved fatal. When atmospheric air containing a mixture of it is breathed, it occasions a violent and almost convulsive cough, attended with much pain in the chest. This cough usually continues to return at intervals for a day or two, and is accompanied with a copious expectoration.
- 2. It is espable of supporting combustion; in many cases even more capable than common air. When a

e fift Trees, then.

burning taper is plunged into it, the flame is diminish. Chap. If. ed, and acquires a very red colour; a great quantity of smoke is emitted, and at the same time the taper consumes much more rapidly than in common air . The facility with which bodies take fire in this gas seems to depend on the ease with which it parts with its oxygen.

4. This gas is neither altered by exposure to light nor to caloric. It passes unaltered through red hot porcelain tubes †.

5. It does not unite readily with water. Schoole Action of found, that after standing 12 hours over water, #ths of the gas were absorbed : the remainder was common air. which no doubt had been contained in the vessel before the operation. Berthollet surrounded several bottles containing it with ice: as soon as the water in these bottles was saturated, the gas became concrete, and sunk to the bottom of the vessels; but the smallest heat made it rise in bubbles, and endeavour to escape in the form of gas 1. Westrumb observed that it became solid when exposed in large vessels to the temperature of 40°; and that then it exhibited a kind of crystallization (. The specific gravity of water saturated with this gas, at the temperature of 43°, is 1.003 ||. From Berthollet's experiments, it appears that a cubic inch of water is capable of absorbing about 1.6 grains (French) of this acid gas. Water impregnated with this gas is usually distinguished by the name of oxymuriatic acid. It has a pale greenish yellow colour, and a suffocating

Foureroy, Ann. de Chim. iv. 231.

^{\$} Jour. de Phys. 3785.

Bertholiet, Ibed. 1785.

[|] Pourcroy, il. 210. f lbid. znavil 382,

odour like the gas; its taste is not acid, but astringent. It is usually prepared by causing the gas to pass thro' a succession of Wolfe's bottles nearly filled with pure water. Light decomposes this acid, as Berthollet discovered, though it has no action on the gas.

- other acids do; and the colour thus destroyed can neither be restored by acids nor alkalies. It has the same effects on yellow wax. If the quantity of vegetable colours to which it is applied be sufficiently great, it is found reduced to the state of common muriatic acid.

 Hence it is evident, that it destroys these colours by communicating oxygen. This property has rendered oxymuriatic acid a very important article in bleaching.
 - y. Oxymuriatic acid is not altered by oxygen gas; but all the simple combustibles are capable of decomposing it.

Action of simple combustibles. When one measure of hydrogen gas is mixed with two measures of oxymuriatic acid, and kept for 24 hours in a phial closed with a ground stopper, and the phial is then opened under water, the whole of the gaseous contents disappear. Hence it is obvious that they act upon each other: the hydrogen absorbs the oxygen of the acid, and is converted into water, while muriatic acid is evolved †. When a mixture of oxymuriatic acid gas and hydrogen gas is made to pass through a red hot porcelain take, a violent detonation takes place 1. By electricity a feeble explosion is produced.

This gas was obtained by a method to be described in the next. Section, and obviously contained a portion of hyperoxymuriatic acid.

[†] Czuikshank's, Nicholson's Journal, 1802, v. 202.

[‡] Fourcroy, ii. 110.

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When melted sulphur is plunged into it, inflammation also takes place, and the sulphur is converted into sulphuric acid ; but cold sulphur, though it is exidized by this gas, does not take fire id it †.

When phosphorus is plunged into this gas, it immediately takes fire, burns with considerable splendour, and is converted into phosphoric acid. This was first described by an anonymous German writer 1, and afterwards by Westrumb, Schmeisser, Fourcroy, and Vanguelin 6.

When charcoal in fine powder is thrown into this gas, heated to about 90°, it also takes fire, according to Westrumb; but this experiment has not succeeded in the hands of other chemists. If we believe Professor Lampadius, the diamond also, when heated to redness, and plunged into oxymuriatic acid gas, burns in it with great splendour; but this experiment has also failed in the hands of other chemists.

Sulphureted, phosphureted, and carbureted hydrogen gas likewise decompose this acid; but none of them, except phosphureted hydrogen, produce spontaneous inflammation with it.

When one measure of carbureted hydrogen, from other or camphor, is mixed with two measures of oxymuriztic gas, and allowed to remain for 24 hours in a phial closed with a ground stopper, the gases mutually decompose each other: water, muriatic acid, carbonic acid, and carbonic oxide, are formed. Accordingly,

Fourcroy, ii. 110.

[†] Westrumb, Ureli's Annals, i. 150. Eng. Transl. See also Vol. I. p. 125, of this Work.

Crell's Bettenge, vol. i. part 3.

Ann. de Cb m. iv. 253.

when water is admitted, the whole is absorbed except about 0.43 of a measure: 0.09 of this residue is absorbed by lime-water; the rest is carbonic oxide. When there is an excess of oxymuriatic acid, the resulting substances are water, muriatic acid, and carbonic oxide. When a mixture of two parts oxymuriatic acid gas, and one part carbureted hydrogen, is fired by electricity, charcoal is deposited, and the gas diminished to 0.6 of a measure; 0.5 of which are absorbed by water, the rest is combustible †.

8. Oxymuriatic acid is not acted upon by either of, the simple incombustibles.

Action on metals

9. Oxymuriatic acid oxidizes all the metals without the assistance of heat. Several of them even take fire as soon as they come into contact with the gas, as Westrumb first discovered. All that is necessary is to throw a quantity of the metal, reduced to a fine powder, into a vessel filled with the gas. The inflammation takes place immediately; the metal is oxidized, while the acid, decomposed and reduced to common muriatic acid, combines with the oxide, and forms a muriate. nic burns in oxymuriatic acid gas with a blue and green flame; bismuth with a lively bluish flame; nickel, with a white flame, bordering on yellow; cobalt, with a white flame, approaching to blue; zinc, with a lively white flame; tin, with a feeble bluish flame; lead, with a sparkling white flame; copper and iron, with a red flame 1. Several of the metallic sulphurets, as cinnabar, realgar, sulphuret of antimony, take fire when thrown in powder into this gas.

^{*} Cruikshanks, Nicholson's Journal, 1302, v. 204.
† Id. Ibid.

‡ Jour. de Phys. xxxvii. 385.

10. When oxymuriatic acid gas and ammoniacal gas are mixed together, a rapid combustion, attended with On any a white flame, instantly takes place; both the gases are decomposed, water is formed, while azotic gas and muriatic acid are evolved . The same phenomena are apparent, though in a smaller degree, when liquid ammomais poured into the acid gas +. The same decompoaition takes place though both the acid and alkali be in a liquid state. If four-fifths of a glass tube be filled with oxymuriatic acid, and the remaining fifth with ammonia, and the tube be then inverted over water, an effervescence ensues, and azotic gas is extricated 1. It was by a similar experiment that Berthollet demonstrated the composition of ammonia.

11. This acid has not hitherto been combined with the alkalies, earths, or metallic oxides; nor have sufficient proofs been adduced that it is capable of combining with these bodies.

12. Oxymuriatic acid gas reddens nitrous gas, and converts it into nitrous acid. It produces no effect upon any of the acids hitherto described, except the sulphuzous and phosphorous, which it converts into sulphurio and phosphoric.

13. When muriatic acid is mixed with nitric acid, Aquategia. the compound has precisely the smell and the qualities of exymuriatic. This mixture of the two acids was formerly called aqua regia; but at present it is usually denominated nitro muriatic acid. It is first mentioned by Isaac Hollandus, and seems to have been known be-

[·] Fourcroy, Ann. de Chim. iv. 255.

⁺ Westrumb, Crell's Assales, i. 161. English Transl.

³ Jour. de l'Ecole Polytecon.

fore the muriatic acid itself. It was prepared by pouting nitric acid on common salt. The nitric acid decomposes the salt, and part of it unites with the muriatic acid thus set at liberty. As soon as these two acids are mixed they begin to act upon each other. The muriatic acid decomposes part of the nitrie, combines with its oxygen, and is thus partly converted into oxymuriatic acid. Hence the suffocating odour of that said which the mixture exhales. The nitrous gas, thus set at liberty, is absorbed by the undecomposed nitrie acid, and converts it into nitrous seid: When these mutual combinations are completed, the action of the two seids on each other ceases. Thus nitro-muristic acid is a mixture or combination of nitrous acid, murisho acid, and exymuristic acid *.

Supposed formation of muriatic acid.

Mr Lambe some time ago † announced, that when ison is acted upon by sulphureted hydrogen gas, a substance is produced which possesses all the properties of exymuriate of iron (oxymuriatic acid combined with iron). In a solution of this gas in distilled water, he digested iron filings, previously purified by repeated washings with distilled water. The bottle was filled with the solution, and corked. The iron was presently acted upon; numerous bubbles arose, which drove the cork out of the bottle; they were strongly inflammable, and probably therefore pure hydrogen gas. The liquor gradually lost its odour of sulphureted hydrogen gas, and after some days smelled very much like stagnant rain-water. As the bubbles ceased to be produced, it recovered its transparency. On evaporating a small

^{*} Fourcroy, ii. 107.

^{\$} Manchester Man. v. 194.

quantity of this solution in a watch-glass to dryness, a bitter deliquescent salt was left behind. On this salt a little sulpourie acid was dropped, and paper moistened with ammonia was held over the glass; white vapours were immediately formed over the glass; and consequently some volatile acid was separated by the sulphuric acid. Mr Lambe evaporated about eight ouncemeasures of the same liquor, and, as before, dropped a little sulphuric acid on the residuum; a strong effervescence was excited, very pungent acid furnes arose, which, from their smell, were readily known to be muriatic. The same truth was established beyond a doubt, by holding a bit of paper, moistened with water, which made the vapours visible in the form of a grey smoke; a distinguishing characteristic, as Bergman has obserwed, of the muriatic acid. When manganese and mereury were dissolved in sulphureted hydrogen gas, the salts formed gave the same unequivocal marks of the presence of muriatic acid.

This experiment of Mr Lambe has been lately repeated with every possible precaution by Vauquelin,
and also by Guyton Morveau and Bouillon Lagrange;
but these chemists did not succeed in obtaining a particle of oxymuriate of iron. We must conclude, then,
that in the experiment of Mr Lambe, muriatic acid
must have insmuated itself into his solution by some
unknown channel. A fact mentioned by Berthollet
may perhaps contribute something to explain this insinuation, and may also suggest a valuable hint towards
the investigation of the real component parts of this ob-

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⁴ Ann. de Chim. Exzvii 191.

stinate acid. He converted a quantity of iron into filings with all possible care. The filings, when washed with water, gave no marks of containing muriatic acid; but after being exposed for some days to the air, they furnished, when again washed, evident traces of the presence of muriatic acid.

Component parts of onymuriation

14. From the action of oxymuristic acid on combustibles, above described, and the compounds produced by it, no doubt can be entertained that it is a compound of muriatic acid and oxygen. This was first demonstrated by Berthollet, by a great number of ingenious and decisive experiments.

He attempted also to ascertain the proportion of its constituent parts. For this purpose he saturated 50 cubic inches (French) of water with oxymuriatic acid, and exposed the liquid for some days to the light of the sun. The oxygen gas disengaged amounted to 15 cubic inches (French) or 8 grains. The muriatic acid which remained in the liquid amounted to 65 grains. Hence it follows, that oxymuriatic acid is composed of 65 parts muriatic acid and 8 parts oxygen, or of about

89 muratic acid

11 oxygen

100

But this method, though exceedingly simple and ingentious, was scarcely susceptible of precision. The analysis of Ghenevix, made in a different way, describes more confidence. This celebrated them are considence.

before by Margraff.

ent of oxymuriatic acid gas to pass through a dilu- Chap. II. polution of potash in water, till it contained an exof acid. He then evaporated to dryness, and prod a saline mass containing all the potash, and the muriatic acid. But when oxymuriatic acid is made ct upon potash in this manner, it divides itself into portions; one portion consists of common muriatic l, the other contains all the oxygen, and is therefore he state of hyperoxymuriatic acid. The potash in dry salt was therefore partly combined with muriaacid, and partly with hyperoxymuriatic acid. Nie of silver precipitates the first of these acids from Hence an easy meiolutions, but not the second. I of ascertaining the relative proportion of these two s in a given quantity of salt presented itself. Acingly Mr Chenevix ascertained, that in 100 parts ne dry salt, 84 consisted of 56.12 parts of potash bined with 27.88 of muriatic acid, and 16 of 3.8 stash combined with 12.2 of hyperoxymuriatic acid. he had ascertained, by a method to be described in next section, that 12:2 parts of hyperoxymuriatic contained 9 parts of oxygen and 3.2 of muriatic Therefore 27.88 + 3.2 = 31.08 muriatic acid; 9 parts of oxygen form 40.08 of oxymuriatic acid. ice it follows, that oxymuriatic acid is composed of

> 77.5 muriatic acid 22.5 oxygen

100

5. Though oxymuriatic acid has hitherto been plaamong acids by chemists, it does not possess a single erty which characterises that class of bodies. Its is not acid but astringent; it does not convert vel'ol. 11.

Not an

getable blues to red, but destroys them; it combines very sparingly with water, and is not capable of neutralizing alkalies, earths, or metallic oxides. It ought therefore to be placed among the oxides rather than the acids. But Scheele, the original discoverer of it, was induced, from the theory which then prevailed, to consider it as merely muriatic acid deprived of phlogisten; and after the mistake was discovered, the theory of Lavoisier respecting the acid principle, which then became fashionable, was considered as a sufficient reason for continuing it in its place among acids in spite of as properties. It must be confessed, however, that such arbitrary arrangements are no small impediments to the progress of the science.

SECT. IV.

OF HYPEROXYMURIATIC ACID.

The existence of this acid was first suspected by Mr Berthollet, though he satisfied himself with little more than strong analogy. It has been lately put out of doubt by the experiments of Mr Chenevix.

Fermation

1. If a solution of potash in six times its weight of water be put into a Wolfe's bottle, and a stream of oxymuriatic acid gas be made to pass through it in the usual way till the potash is saturated, crystals in the form of fine white scales are deposited in considerable quantity. These crystals have received the name of hyperoxygenized muriate of potasb. They possess very

purious and important properties. If the liquid from which this sait is deposited be evaporated to dryness, another salt will be obtained, composed of muriatic acid and potash. These facts were all discovered by Berthollet. He concluded from them, that the oxymuriatic acid had been decomposed during the process; that one portion of it lost the whole of its oxygen, and was reduced to the state of muriatic acid, while another portion combined with an additional dose of oxygen, and was converted into byperoxymuriatic acid. Hence the appellation of the salt which contained this last acid.

2. This theory of Berthollet was very plausible; and Opinion of it induced the greater number of chemists to believe that the substance to which hyperoxymuriate of potash owes its peculiar properties, differs from oxymuriatic aacid by containing an additional dose of oxygen. But the opinion remained destitute of sufficient proof, till Mr Chenevix published his important dissertation on the subject in 1802. Mr Chenevix exposed 100 grains of hyperoxymuriate of potash to the heat of a lamp; it lost 2.5 parts of its weight, which he ascertained to be water. When heated to redness, a violent efferves- Confirmed cence took place, and 112.5 cubic inches of oxygen by Chene gas, or 38-3 grains, were extricated. The salt which remained in the retort amounted to 53.5 grains, and five grains had been volatilized during the process . Hence it follows, that hyperoxymuriate of potash is composed of

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It had been long known that this saline residue is a compound of porash and muristic soid.

ACID SUPPORTERS.

Book II. Division II. 2.5 water

38.3 oxygen

58.5 muriate of potash

99.3

But Mr Chenevix ascertained, that the mariatic acid remaining in this saline residue amounted to 20 grains. Therefore 38.3 parts of oxygen, and 20 parts of muritic acid, constitute 58.3 parts of the acid which exists in hyperoxymuriate of potash. That acid of course is composed of about.....66 oxygen

34 muriatic acid

100

The acid which constitutes a part of hyperoxymuriate of potash contains, we see, nearly nine times as much oxygen as exists in oxymuriatic acid. Thus the theory of Berthollet is confirmed in the fullest manner. We see that muriatic acid combines with two doses of oxygen; with the first dose it constitutes oxymuriatic acid, with the second the acid that exists in hyperoxymuriate of potash, which has been called opperoxymuriatic acid.

Oxymuriatic acid contains.....0.22 oxygen

Hyperoxymuriatic acid.....0.66

Hence it follows, that

Muriatic Oxymur. acid. Oxygen. acid.

1.00 + 0.29 = 1.29 Oxygen. Hyperoxymeriscic scid.

1.50 + 1.05 = 5.01

Cannot be obtained s.parate.

3. But though the peculiar nature of hyperoxymuriatic acid has been thus demonstrated, all attempts to procure it in a separate state have hitherto failed. Its properties therefore are but imperfectly known. From the amazing energy with which hyperoxymuriate of

potash acts upon combustible bodies, it is obvious that Chap. II. it possesses in perfection the property of supporting combustion. It combines also with alkalies, earths, and metallic oxides, and forms salts of a very peculiar nature, which we shall afterwards examine. Mr Chenevix has rendered it probable, that it converts vegetable blues into red. It cannot be doubted that it combines with water. Several phenomena indicate, that when pure it assumes the gaseous state.

4. When sulphuric acid is poured upon hyperoxy- Action & genized muriate of potash, a violent decrepitation takes place, and sometimes a flash of light is visible. The sulphuric acid in this case combines with the potash, and disengages the hyperoxymuriatic acid. This last acid rises in the state of a heavy vapour of a greenish yellow colour; its smell has some resemblance to that of nitrous gas, but peculiarly fetid: it is compared by Mr Chenevix to the odour emitted by brick kilns, mixed with that of nitrous gas. At the bottom of this vapour is a bright orange-coloured liquid, consisting of the sulphuric acid, the potash, and a portion of the hyperoxymuriatic acid. But the acid thus separated is not pure, being partly decomposed by the process, and being mixed with a portion of sulphuric acid. If we attempt to separate the hyperoxymuriatic acid by distillation, the moment the mixture is heated to about 120°, a violent explosion takes place, which breaks the vessels in pieces. This seems to be owing to the rapidity with which the acid is decomposed by a moderate heat *. Nitric acid produces nearly the same effects as the sulphuric.

[#] Hoyle, Manchester Memoirs, v. 220.—Chenevix, Phil. Trans. 1802.

Division II.

Of muriatic acid.

5. When muriatic acid is poured upon the salt, a violent effervescence takes place, and gas is emitted abundantly, which has the smell and colour of oxymnriatic acid, but is much more rapidly absorbed by water. This process was first pointed out by Mr Cruikshanks. He obtained in this manner the gas which he employed in his experiments on carbonic oxide. When two measures of hydrogen gas are mixed with 2.3 measures of gas procured in this manner, Mr Cruikshanks found that the mixture explodes feebly with the electric spark, and is totally converted into water and muriatic acid. But two measures of hydrogen gas require one measure of oxygen gas to convert them into water. Hence Mr Cruikshanks infers, that 2.3 parts of the gas procured according to his process, contain one part of oxygen and 1.3 of muriatic acid; which gives as its component parts.....56.5 muriatic acid

100.0

43.5 oxygen

This shows us that the gas extricated from the salt by muriatic acid is not pure hyperoxymuriatic acid. Mr Chenevix supposes that it is a mixture of that acid and oxymuriatic acid gas: a portion of the first acid being decomposed by the muriatic acid, with which it comes in contact at the moment of its disengagement. But there are some reasons to question this opinion. When water, impregnated with oxymuriatic acid gas, obtained by Cruikshank's method, is mixed with liquid ammonia, scarcely any gas is extricated. The two bodies combine and form a salt.

^{*} Nicholson's Journal, 1802, v. 206.

Much still remains to be done before the properties this interesting acid be fully developed. Mr Chewix has ascertained, that it is always formed when nimuriatic acid is boiled upon platinum, and likewise en oxide of titanium is precipitated by potash from trintic seid.

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SECT. V.

OF ARSENIC ACID.

ARSENIC, as we have seen formerly, is capable of combining with two doses of oxygen; and both the impounds which it forms with that body possess acid poperties. The first, distinguished by Fourcroy by the one of arsenious acid, but more usually denominated bite oxide of arsenic, has been already described. The scond, called arsenic acid, was discovered by Scheele 1775 t, and its most remarkable properties investited. Pelletier afterwards published a valuable disestation on it. And more lately its properties and onstituent parts have been more completely investigad by Proust and Bucholz 1.

1. Arsenic acid is usually prepared by the process Preparapointed out by Scheele. Three parts of white oxide of rsenic are dissolved in seven parts of muriatic acid, and the solution is mixed with five parts of nitric acid, and distilled to dryness. What remains is arsenic acid. But his method has been considerably improved by Busholz. His method is as follows. Mix in a retort one

[•] See Vol. L. p. 347.

⁺ Scheele, i. 129.

¹ Jour. de Chim. iv. 5.

part of muriatic acid of the specific gravity 1.2, four parts of the white oxide of arsenic, and 12 parts of nitric acid of the specific gravity 1.25. Boil the mixture till the oxide disappear and nitrous gas ceases to be disengaged. Then evaporate to dryness, and expose the mass for a few minutes to a low red heat. What remains after this is solid arsenic acid.

Properties.

2. Arsenic acid, thus prepared, is a white solid man, nearly tasteless. Its specific gravity is 3.391. It is very fixed. When heated strongly, it melts and remains transparent, and is converted into a glass, which acts powerfully on the vessel in which the experiment is performed. This glass attracts moisture from the air. When the heat is very strong, the acid gives out a little oxygen gas, and is at the same time partly converted into white oxide.

Action of water.

3. It dissolves very slowly in six parts of cold water; but two parts of boiling water dissolve it almost instantly, and it remains in a state of solution even tho a considerable portion of that water be evaporated. With half its weight of water it has a syrupy consistency; and by farther evaporation it deposites crystals in grains *. Its taste, when liquid, is acid, caustic, and metallic.

Of simple hodies.

- 4. Oxygen has no action whatever on this acid; neither is it affected by exposure to the open air.
- 5. The simple combustibles decompose it by the assistance of heat: And in these cases, as we learn from the experiments of Scheele and of the Dijon Academicians, combustion sometimes takes place; a proof that arsenic acid is a supporter of combustion.

Bucholz, Jour. de Chim. iv. 5.

6. The simple incombustibles do not appear to pro- Chap. II. duce any alteration on it.

- 7. Several of the metals decompose it when assisted It does not act upon gold, platinum, silver, mercury. It oxidizes copper, iron, lead, tin, zinc, bismuth, antimony, cobalt, nickel, manganese, and arsenic, and in a strong heat mercury and silver.
- 8. It combines with alkalies, earths, and several of the metallic oxides, and forms compounds which have . been called arseniates.
- 9. From the way in which this acid is formed, it Composicannot be doubted that it is composed of the white oxide of arsenic and oxygen. Nor is it very difficult to ascertain the proportion of these constituent parts; for the oxygen must be equivalent to the increase of weight which the oxide experiences during its acidification. From the experiments of Proust we learn, that this augmentation of weight amounts to 0.15 parts: Bucholz found it 0.16; a degree of coincidence as great as can be expected in experiments of that delicate nature: Hence it follows that arsenic acid is composed of about

86.5 white oxide of arsenic

13.5 oxygen

100.0

But the oxide of arsenic contains nearly 0.25 of oxygen. This gives us arsenic acid composed of about

65 arsenic

35 oxygen

100

It is only the second dose of oxygen, amounting to 13.5 parts, which gives this acid the property of sup-

porting combustion, and of emitting oxygen gas when strongly heated.

SECT. VI.

OF TUNGSTIC ACID.

History.

THE substance called tungstic acid by Scheele and Bergman was discovered by Scheele in 1781. This philosopher obtained it from tungstate of lime by treating it with nitric acid and ammonia alternately. The acid dissolves the lime, and the ammonia combines with the tungstic acid. The ammoniacal solution, when saturated with nitric or muriatic acid, deposites a white powder, which is the tungstic acid of Scheele.

This powder has an acid taste, it reddens vegetable blues, and is soluble in 20 parts of boiling water. The De Luyarts have demonstrated, that this pretended acid is a compound of yellow oxide of tungsten, the alkali employed to dissolve it, and the acid used to precipitate it. Thus, when prepared according to the above described process, it is a compound of yellow oxide, ammonia, and nitric acid. Their conclusions have been more lately confirmed by the experiments of Vauquelia and Hecht. This substance must therefore be erased from the class of acids, and placed among the salts.

The real acid of tungsten is a yellow powder; the method of procuring which, and its properties, have been already described under the denomination of yel-

oxide of tungsten . It ought rather, as Vauque- Chap II. and Hecht have properly remarked, to be classed mong the oxides than the acids; for it is insoluble in enter, tasteless, and has no effect on vegetable blues. lagrees with the acids indeed in the property of comming with alkalies and earths, and perhaps also with me metallic oxides, and forming with them salts, bich have been denominated tungstates; but several mer metallic oxides, those of lead, silver, and gold, instance, possess the same property. These oxides erefore may be called acids with as much propriety as yellow oxide of tungsten.

SECT. VII.

OF MOLYBDIC AGID.

MOLYBDIC acid was discovered by Scheele in 1778, History. mering his experiments on the sulphuret of molybdemm, and its most remarkable properties ascertained. searcely any farther addition was made to our knowedge of it, till Mr Hatchett published his dissertation the molybdate of lead, in the Philosophical Tranections for 1796. That ingenious chemist examined. ith his usual precision, such of its properties as were connected with this subject. In the summer of 1805. elaborate dissertation on molybdenum was publishin the 4th volume of Gehlen's Journal, by Bucholz,

who did not neglect to ascertain the constituents and the mode of procuring this acid. To Scheele, Hatchet, and Bucholz, we owe most of the facts respecting molybdic acid at present known.

From the experiments of Bucholz it appears, that two of the combinations of molybdenum and oxygen are soluble in water; namely, the blue oxide and the peroxide. If the first be capable of neutralizing alkalies, it will belong to the class of acids, and may be distinguished by the name of molybdous acid; but it acid properties have not been ascertained. The peroxide, however, neutralizes alkalies. It has been always distinguished by the name of molybdic acid.

Prepara-

It is usually prepared from molybdena, or native sulphuret of molybdenum, by the process pointed out by Scheele. Reduce the mineral to powder, and distil off it nitric acid, or rather a mixture of nitric and muriatic acids, till the whole is converted into a white mass. Edulcorate this mass with water to carry off the sulphuric acid formed, and the remains of the other acids. It is now molydic acid tolerably pure.

Another method has been lately pointed out by Bucholz. Reduce the mineral to a fine powder, and expose it to heat in an open crucible, stirring it with an iron rod till the whole assumes an ash grey colour. The heat is to be at first a strong red, but gradually lowered as the roasting advances, to prevent the powder from cohering, which would render the completion of the process very difficult. By this roasting the sulphur is dissipated, and a considerable portion of the metal acidified. Reduce the mass to powder, and digest it a sufficient time in water holding soda or ammonia in solution; the molybdic acid is taken up, and combines

with the alkali, while the impurities remain behind. Chap. II. Let the solution remain corked up till it has become clear, decant it off from the sediment if any thing has subsided, and pour into it some muriatic acid. The molybdic acid precipitates in the state of a fine white powder, and may be easily separated and edulcorated *.

Molybdic acid, thus prepared, is a white powder, the Properties specific gravity of which is 3.460. When heated in a close vessel it melts and crystallizes; but in an open vessel it sublimes in a white smoke, which attaches itself to cold bodies, and assumes the form of brilliant yellow scales.

It is soluble in 960 parts of boiling water. The solution is pale yellow; it has no taste, but reddens litmus paper. The molybdic acid is precipitated from this solution by sulphuric, nitric, and muriatic acids +.

Molybdic acid is not affected by oxygen gas; but it is decomposed by sulphur and charcoal, and several of the metals. When heated with the protoxide of molybdenum, the mixture is converted into blue oxide 1.

It combines with alkalies, earths, and several metallic oxides, and forms salts known by the name of molybdates.

Sulphuric acid dissolves molybdic acid when assisted The solution is colourless while hot; but when cold it assumes a deep blue colour, which is heightened by saturating the solution with soda. When this sulphuric acid solution is heated strongly, the sulphuric acid is evaporated, and molybdic acid remains. Muriatic acid also dissolves it. The solution is of a

[#] Gehlen's Jour. iv. 604.

⁺ Hatchett, Phil. Irens. Ixxxvi. 323.

² Bucholz, Gehlen, iv. 626.

pale yellowish green colour; but it becomes blue when saturated with potash. Nitric acid does not dissolut this acid .

This acid, when combined with potash, forms a colourless salt.

Mixed with filings of tin and muriatic acid, it immediately becomes blue, and precipitates flakes of the same colour, which disappear after some time, if an excess of muriatic acid has been added, and the liquor assumes a brownish colour.

With the solution of nitrate of lead it forms a white precipitate, soluble in nitric acid.

When mixed with a little alcohol and nitric acid, it does not change its colour.

With a solution of nitrate of mercury, or of nitrate of silver, it gives a white flaky precipitate.

With the nitrate of copper it forms a greenish precipitate.

With solutions of sulphate of zinc, muriate of bismuth, muriate of antimony, nitrate of nickel, muriates of gold and platinum, it produces white precipitates when these solutions do not contain an excess of acid.

When melted with borax, it gives it a bluish colour.

Paper dipt in this acid becomes in the sun of a beautiful blue colour +.

From the experiments of Bucholz we learn, that when 100 grains of molybdenum are digested with nitric acid till they are converted into molybdic acid, and then dried, they now weigh about 149 grains. Hence it follows, that molybdic acid is composed of about 100

B Hatchett, Phil. Trans. lxxxvis 323. + Vauquelin, Phil. May i. 282.

marts metal and 49 oxygen; or very nearly two parts Chap. IL metal to one oxygen; or per cent. of about

67 molybdenum

33 oxygen

100

SECT. VIII.

OF CHROMIC ACID

CHROMIC ACID, discovered lately by Vauquelin, has Preparaonly been found, in any quantity, in the red lead ore of Siberia, and in chromate of iron. Its properties have been investigated by Vauquelin and Mussin Paschkin.

It may be obtained by boiling the red lead ore with carbonate of soda, decanting off the fluid solution, and saturating it with one of the mineral acids; a red powder precipitates, which is chromic acid.

Chromic acid, thus obtained, is a red or orange-yel- Properties. low powder, of an acrid and strongly metallic taste. It is soluble in water, and crystallizes in the form of elongated prisms of a ruby colour.

When heated it gives out oxygen gas, and is converted into green oxide of chromium.

When mixed with filings of tin and the muriatic acid. it becomes at first yellowish brown, and afterwards ussumes a beautiful green colour.

[@] Gchlen's Jow. iv. 618.

When mixed with a little alcohol and nitric acid, it immediately assumes a bluish green colour, which preserves the same shade even after desiccation. Ether alone gives it the same colour.

With a solution of nitrate of mercury, it gives a precipitate of a dark cinnabar colour.

With a solution of nitrate of silver, it gives a precipitate which, the moment it is formed, appears of a beautiful carmine colour, but becomes purple by exposure to the light. This combination, exposed to the heat of the blowpipe, melts before the charcoal is inflamed. It assumes a blackish and metallic appearance. If it be then pulverised, the powder is still purple; but after the blue flame of the lamp is brought in contact with this matter, it assumes a green colour, and the silver appears in globules disseminated throughout its substance.

With nitrate of copper, it gives a chesnut red precipitate.

With the solutions of sulphate of zinc, muriate of bismuth, muriate of antimony, nitrate of nickel, and muriate of platinum, it produces yellowish precipitates when these solutions do not contain excess of acid. With muriate of gold it produces a greenish precipitate.

When melted with borax or glass, it communicates to them a beautiful emerald green colour.

Paper impregnated with chromic acid assumes in the light a greenish colour.

When mixed with muriatic acid, the mixture is capable of dissolving gold like aqua regia: when this mixture of the two acids is distilled, oxymuriatic acid is disengaged, and the liquor assumes a very beautiful green colour.

Sulphuric acid, while cold, produces no effect upon but when warmed, it makes it assume a bluish reen colour, probably by favouring the disengagement of axygen.

When this acid is heated along with charcoal, it is educed to the metal called chromium.

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SECT. IX.

F COLUMBIC ACID.

THIS metallic acid has hitherto been detected in one Preparapecimen only, in which it is combined with the oxide tion. iron. Mr Hatchett, to whom we owe the discovery of its peculiar properties, separated it from the iron by basing it with potash. The alkali combined with a portion of the acid which was separated by water. Muristic acid dissolved the oxide of iron thus deprived of its acid; and the ore, after this treatment, yielded an additional dose of acid when fused with potash. By ree peating this process, the whole of the acid was combined with potash: the solution was limpid. Nitrie acid precipitated the columbic acid from the potash in the state of white flakes.

- 1. The acid thus obtained is of a pure white colour, Properties. and not remarkably heavy. It has scarcely any taste. It is insoluble in water, but gives a red colour to paper stained with litmus. When exposed to a strong heat, it does not melt, but loses its lustre.
- 2. Sulphuric acid dissolves it, and forms a transparent co.ourless solution: But when the aquid is diluted

with water, it becomes milky; a white precipitate falls, which becomes lavender blue; and when dry, brownish grey. It is semitransparent, and breaks with a vitreous fracture. This substance is a compound of salphuric and columbic acids.

- 3. Nitric acid does not dissolve this acid; nor does it produce any change on its properties.
- 4. Muriatic acid dissolves it when assisted by heat. The solution does not become muddy when diluted with water. When evaporated to dryness, it leaves a pale yellow substance difficultly soluble in muriatic acid.
- 5. Potash and soda combine readily with columbic acid. With potash this acid forms a glittering scaly salt like boracic acid. It contains a slight excess of alkali; has an acrid disagreeable flavour; does not dissolve readily in cold water; but when dissolved, the solution is perfect and permanent. Nitric acid precipitates the columbic acid. But columbic acid is capable of expelling carbonic acid from potash.

The other properties of this acid remain to be investigated.

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CB

EL

[#] Hatchett, Phil. Tress. 1802.

CLASS III.

COMBUSTIBLE ACIDS.

TE acids belonging to this class were formerly distinished by the appellation of vegetable and animal acids, cause almost the whole of them are procured from the amal and vegetable kingdom. They differ essentially m the other two classes of acids in several particulars.

1. If they be combined with potash and distilled, Properties. er are completely decomposed, charcoal is usually eolved, and a considerable quantity of heavy inflamsable air extricated; whereas no combustible substance a be procured by exposing the other acids to heat *.

2. All of them contain at least two simple combus-The substances as a base; whereas the others never conin more than one. These two substances are always wrbon and bydrogen. Some of these acids contain likewise avote, over and above the two simple combusables. Oxygen also usually enters into their composition in considerable quantity, but not perhaps always. To them, therefore, the theory of Lavoisier, respecting

Sulphurous and phosphorous acids excepted, which entit sulphur and phosphureted hydrogen when exposed to heat.

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the necessity of oxygen as the acidifying principle, does not strictly apply.

- 3. They do not seem capable of combining with different proportions of oxygen. Whenever the quantity of oxygen is changed, the proportion of the other constituents changes also. Of course the termination of their names ought not to indicate the proportion of oxygen which they contain; but should, if possible, be to dependent of that proportion altogether.
- 4. They are decomposed by the action of the monpowerful acid supporters, and converted either into other combustible acids; or each of their constitues is saturated with oxygen, and converted into oxide miacid products.

Divisible into four orders.

5. The combustible acids may be very conveniently subdivided into four orders; which are distinguished from each other by the following properties:

The acids belonging to the first order are crystallizable, and they may be volatilized by heat without usdergoing decomposition.

Those belonging to the second order are likewiss crystallizable, but they cannot be volatilized without decomposition.

The French chemists who formed the new nomenciature were miled here. They made some of the names of the combustible acids at in m, as if they were saturated with oxygen; and others in m, at if they were capable of combining with an additional dose. The fact is, that none of them are, strictly speaking, saturated with oxygen; for it of them are capable of combining with more. But then they cannot combine with more without being totally decomposed, and converted my water, carbonic acid, &cc. I have adopted the rule proposed by Mr Chanevix, which is calculated to avoid all ambiguity; the names of all the combostible acids are made to end in m, and this without reference to themy.

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the fourth order are placed three acids, which, singularity of their properties ought to be secon the rest. For want of a better name we inguish them by the term colorific.

Howing Table contains the names and commets of all the acids belonging to each of these far as they have been ascertained.

DER I. Crystallizable. Volatilizable.

illion.	Constituents.
tic toic teic tanic toxylic tohoric	Carbon, hydrogen, oxygen.

Table of the constituent parts.

II. Crystallizable. Not volatilizable.

litic taric te te	Carbon, bydrogen, oxygen.
i i	Carbon, hydrogen, azote, oxygen.

ORDER III. Not crystallizable.

ric nic	Carbon, hydrogen, oxygen.
- T	

ORDER IV. Colorific.

.]	Names.	Constituents.	
	Gallic	Carbon, hydrogen, exygen.	
_	Prussic	Carbon, hydrogen, azote.	
•	3. Sulphureted hydrogen	Sulphur, hydrogen.	

We shall consider the properties of the acids be to the three first orders in the following Sections fourth order, for reasons to be assigned hereafts occupy a separate Chapter.

SECT. I.

OF ACETIC ACID.

This acid is employed in three different states have been distinguished from each other by names. When first prepared, it is called when purified by distillation, it assumes the distilled vinegar, usually called acetous acid by characteristic acid. It will be necessary to describe each states separately.

Vinegar.

1. Vinegar was known many ages before to covery of any other acid, those only excepted exist ready formed in vegetables. It is mention Moses, and indeed seems to have been in comme

mong the Israelites and other eastern nations at a very Chap. It. arly period. It is prepared from wine, from beer, ale, ather similar liquids. These are apt, as every one www, to turn sour, unless they be kept very well cork-Now sour wine or beer is precisely the same with Aregar.

Boerhaave describes the following method of making anegar, which is said to be still practised in different Laces.

Take two large oaken vats or hogsheads, and in each Preparaf these place a wooden grate or hurdle at the distance a foot from the bottom. Set the vessel upright, and on the grate place a moderately close layer of green wigs of fresh cuttings of the vine. Then full up the vessel with the footstalks of grapes, commonly called the rape, to the top of the vessel, which must be left quite open.

Having thus prepared the two vessels, pour into them the wine to be converted into vinegar, so as to fill one of them quite up, and the other but half full. Leave them thus for twenty-four hours, and then fill up the half-filled vessel with liquor from that which is quite full. Four-and-twenty hours afterwards repeat the same operation; and thus go on, keeping the vessels alternately full and half full during every twenty-four hours till the vinegar be made. On the second or third day there will arise, in the half-filled vessel, a fermentative motion, accompanied with a sensible heat, which will gradually increase from day to day. On the contrary, the fermenting motion is almost imperceptible in the full vessel; and as the two vessels are alternately full and half full, the fermentation is by that means, in

some measure, interrupted, and is only renewed every other day in each vessel.

When this motion appears to have entirely cessed, even in the half-filled vessel, it is a sign that the fermentation is finished; and therefore the vinegar is then to be put into casks close stopped, and kept in a couplace. The process in summer usually lasts fifteen days; in winter it lasts longer.

All that is necessary to convert wine or beer into vinegar is the contact of the external air, a temperature of 80°, and the presence of some substance to act as a ferment. But the theory of this operation belongs to the Second Part of this Work; our business at present is, not to investigate the method of making vinegar, but to examine the properties of acetic acid.

Properties.

Vinegar is a liquid of a reddish or yellowish colour, a pleasant sour taste, and an agreeable odour. Its specific gravity varies from 1.0135 to 1.0251, and it differs also in its other properties according to the liquid from which it has been procured. It is very subject to decomposition; but Scheele discovered, that if it be made to boil for a few moments, it may be kept afterwards for a long time without alteration. Besides acetic acid and water, vinegar contains several other ingredient, such as mucilage, tartar, a colouring matter, and often also two or more vegetable acids. When distilled at a

An account of the mode of making vinegar in France, almost precisely the same with that given in the text, was published in the Philosophical Transactions for 1670, vol. v. p. 2002. The methods at present practised in France are described in the 3d Volume of the Element de Chymic, published by the Dijon Academy, p. 6. Many still follow the process described in the text; others a different one,

temperature not exceeding that of boiling water, till a- Chap II. bout two-thirds, or five-sixths at most, of it have passed over, all these impurities are left behind, and the product is pure acid, diluted with water. The residuum is still an acid liquid, and often lets fall crystals of

2. The acid thus obtained is a liquid as transparent Acctous and colourless as water, of a strong acid taste and an agreeable odour, somewhat different from that of vince gar. In this state it is usually called acetous acid or distilled vinegar.

It may be preserved without alteration in close vessels. When exposed to a moderate heat, it evaporates completely and without undergoing any charge in its properties. When exposed to the action of cold, part of it congeals. The frozen portion, which consists almost entirely of water, may be easily separated; and by this method the acid may be obtained in a high degree of concentration. The more concentrated the acid is, the greater is the cold necessary to produce congelation. Mr Lowitz has ascertained that the acid itself, how much soever it be concentrated, crystallizes or congeals at the temperature of -22°.

3. When acetate of copper, reduced to powder, is put Preparainto a retort and distilled, there comes over a liquid at coucacid. first nearly colourless and almost insipid, and afterwards a highly concentrated acid. The distillation is to be continued till the bottom of the retort is red hot. What remains in it then is only a powder of the colour of copper. The acid product, which should be received. in a vessel by itself, is tinged green by a little copper which passes along with it; but when distilled over again in a gentle heat, it is obtained perfectly colourless

and transparent. The acid thus obtained is exceeding. ly pungent and concentrated. It was formerly distinguished by the names of radical vinegar and vinegar of Venus.

This process was known to the alchymists, and had often been repeated by chemical philosophers; but the product was considered as merely highly concentrated acetous acid, till Berthollet published his experiments on it in 1785 *. That skilful philosopher affirmed, the it differs from acetous acid in taste and smell, in its affinities for other bodies, and in the compounds which it forms with them. When it is obtained from acetate of copper by distillation, the powder which remains in the retort consists chiefly of copper in the metallic state. Hence he supposed, that the acetic acid gave out phlogiston to the copper, and received from it oxygen. When the existence of phlogiston was disproved, this theory was a little altered. It was concluded, that during the distillation, the acetic acid deprived the oxide of copper of its oxygen, and combined with it; and that radical vinegar therefore is acetous acid combined with a new dose of oxygen. For this reason it received the name of acctic acid.

Supposed different from acc-

This theory was generally admitted by chemists, till Mr Adet published his experiments on acetic acid in 1797†. He observed, that when acetate of copper is distilled, the products are not only acetic acid and water, but likewise carbonic acid and carbureted hydrogen gas; and that the residuum consists of copper and charcoal. He found that acetous acid did not absorb

^{*} M.w. Var. 1783.

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bygen when distilled off black oxide of manganese, and that acette and acetous acids have exactly the same feet upon metals. From these and some other similar periments, he concluded that acetic and acetous acids not differ from each other except in concentration.

This conclusion was considered as hasty, as he did ot attempt to account for the very striking difference the taste and smell of these acids, as he had neither repeated nor refuted the experiments of Berthodet, which the opinion of the difference between these wo acids was founded. The conclusion of Adet was poposed in 1798 by Chaptal, who published a set of exeriments on the same subject *. This philosopher adeavoured to demonstrate, as Berthollet had done before him, that acetic and acetous acids, even when of he same strength, possess different properties, and have fifferent effects on other bodies. From the effect of alphuric acid upon each, and from the quantity of charwhich remains in the retort when equal quantities each saturated with potash are distilled, he concluded hat acrous acid contains a smaller proportion of caron than acetic acid. And this he considered as the cause of the difference in their properties.

In 1800 a new set of experiments on the same sublect was published by Mr Dabit of Nantes †. This
hemist endeavoured to prove that acetic acid contains
greater proportion of oxygen than acctous. When he
distilled a mixture of acetate of potash and sulphuric
seid, the product was acetic acid; but when he substisted muriatic acid for sulphuric, he obtained acetous

^{* .}fan, de Chim. xxviii. 113.

acid; but when to this last mixture a little black oxide of manganese was added, he assures us that he obtained acetic acid. And in none of these cases was there any carbonic acid emitted till towards the end of the distillation.

Proved to be the same.

The experiments of Darracq have at last finally seltled this dispute, by demonstrating that acetous and acetic acids differ merely in concentration. This able chemist repeated the experiments of Adet, which he found perfectly accurate, and performed several new ones; all of which left no doubt that acetous and acetic acids contain precisely the same proportion of oxygen. He next tried the opinion of Chaptal respecting the proportion of carbon in each. When equal quantities of acetic and acetous acids are combined with potash or soda, the dry salts distilled leave the same quantity of charcoal, and yield the same quantity of other products. When reduced to the same specific gravity, the two acids form precisely the same salts with all alkaline and earthy bases. Finally, acetous acid gradually becomes acetic acid when distilled repeatedly off dry muriate of lime; and in this case no gaseous body whatever is evolved. From these facts, it is impossible to entertain a doubt that the two substances are essentially the same, and that these apparent differences are owing to the great quantity of water with which acetous seid is diluted, and the mucilaginous matter which it still retains .

The same results were obtained also by Proust, who had drawn the same conclusions before he came ac-

Darracq, Ann. de Chim. al . 264.

mainted with Darracq's paper *. Henceforth the term ectous acid must either be wholly dropt by chemists, r employed in a sense different from what was formeraffixed to it.

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4. This acid is transparent and colourless like water. Properties. It has a peculiar aromatic smell when in the state of ecetous acid; but concentrated acetic acid, when procuted in the usual way, has an empyreumatic odour mixed with the natural smell of vinegar, owing to a small portion of oil formed during the process +.

Another method of procuring this acid in a concentrated state, has been proposed by different chemists, and brought to a state of perfection by Lowitz of Pepersburgh. Distil a mixture of three parts of acetate of potash and four parts of sulphuric acid, till the aceacid has come over into the receiver. To separate the sulphuric acid with which it is in some measure contaminated, rectify the liquid by distilling it off a The acid that comes portion of acetate of barytes. over crystallizes in the receiver.

The specific gravity of distilled vinegar varies from 1.007 to 1.0095; but radical vinegar is much more concentrated, its specific gravity being as high as 1.080 1. In that state it is extremely pungent and acrid; and when it is applied to the skin, it reddens and corrodes it in a very short time. It is exceedingly volatile; and when heated in the open air, takes fire so readily, that one would be tempted to suspect the presence of ether in it. It unites with water in any pro-

[†] Ann, de Clim, untvii Tat. " Jose de Pbye, lvi. 210. ! Rachter. Gehlen's Jour. iv. 22.

portion; and when concentrated, the mixture evolves a good deal of heat.

5. Courtenvaux had long ago observed, that the last portions of acetic acid, which come over during the distillation, were susceptible of crystallizing in a moderate cold. Lowitz proposed a very ingenious method to obtain this acid in the state of crystals. He makes distilled vinegar into a paste with well burnt charcoal, and exposes the mixture to a heat not above 212°. The watery part is driven off, and the acid remains. A stronger heat drives off the acid itself in a very concentrated state. By repeating this process it may be obtained in crystals. The process of that chemist, detailed above, is more recent, and in every respect preferable.

Action of simple bodies-

- 6. Neither oxygen gas nor air have any very marked action on this acid: nor do the simple combustibles or incombustibles alter it at the common temperature of the atmosphere.
- 7. It is capable of oxidizing iron, zinc, copper, nickel, tin. It does not act upon gold, silver, platinum, mercury, bismuth, cobalt, antimony, arsenic. Its action on tellurium, tungsten, molybdenum, uranium, titanium, and chromium, has not been tried.
- 8. It combines with alkalies, earths, and metallic oxides, and forms compounds known by the name of acetates.
- 9. It is decomposed by sulphuric and nitric acids. The action of the other acids on it has scarcely been examined. It dissolves boracic acid, and absorbs carbonic acid.
 - 10. It has the property of combining with a great

sumber of vegetable bodies, such as bils, mucilage, and Fornatics.

11. When nitric acid is made to act on this acid, it Compos converts it into water and earbonic acid. When acetic scid, combined with a fixed alkali or earth, is exposed to a strong heat, it is almost completely decomposed; water, carbonic acid, and carbureted hydrogen gas, are emitted, and the base remains mixed with a quantity of charcoal. From these facts it follows, that it is composed of carbon, hydrogen, and oxygen, into which all these products may be ultimately resolved. But the proportion of these ingredients has not been ascertained with precision.

By distilling 7680 grains of acetate of potash, Dr Higgens obtained the following products ":

Potash 3862'994 grains Carbonic acid gas 1473'564 Carbureted hydrogen gas 1047'6018 Residuum, consisting of charcoal 78.0000 Oil 180.0000 Water 340'0000 Deficiency †726.9402

This deficiency Dr Higgins found to be owing to a quantity of water and oil which is carried off by the clastic fluids, and afterwards deposited by them. He calculated it, in the present case, at 700 grains of water

" Higgins on Actour And, p. 26.

[†] For 29 1 grains of oxygen gas had also disappeared from the zir of the ventela-

and 26.9402 grains of oil. Now, since acetate of potaticis composed of acetic acid and potash, and since the whole of the potash remained unaltered, it follows, that the acid was converted into carbonic acid gas, exabored hydrogen gas, carbon, oil, and water; all of which are composed of oxygen, hydrogen, and carbon,

Now 1473'564 grains of carbonic acid gas are composed of 1060'966 grains of oxygen and 415'598 grains of carbon.

a comparison of the experiments of Dr Higgins and Lavoisier, may be supposed to consist of about 714'6005 grains of carbon and 333'0010 of hydrogen.

200.9402 grains of oil contain 103.4828 grains of carbon and 43.4574 grains of hydrogen.

1040 grains of water contain \$84 grains of oxygen and 156 grains of by drogen.

Therefore 3617 006 grains of acetic acid are composed of 1944.906 —29.1 = 1915.866 grains of oxygen, 532.4564 grains of bydrogen, and 1368.6816 grains of carbon. Consequently 100 parts of acetic acid are composed of

> 50.19 oxygen 13.94 hydrogen 35.87 carbon

100.00

These numbers can only be considered as very imperfect approximations to the truth; for the object of Dr Higgins was not to ascertain the proportions of the ingredients which compose acetic acid, and therefore has experiments were not conducted with that rigid accuracy which would have been necessary for that put Ose. Mr Proust endeavoured to prove that azote like- Chap. II. rise is a component part of acetic acid; an opinion which had been entertained by Lavoisier. But when Frommsdorf repeated the experiments of Proust, which on sisted in distilling different acetates, he obtained o traces either of ammonia or of prussic acid, as had een announced by the chemist of Madrid. Hence he concludes that acetic acid contains no azote, and that Proust's experiments must have been made upon imare seid ".

SECT. II.

OF BENZOIC ACID.

BENZOIN, or BENJAMIN as it is sometimes called, is a History. kind of resin brought from the East Indies; obtained, according to Dr Dryander, from the styrax benzoe, a tree which grows in the island of Sumatra. This substance consists partly of a peculiar acid, described as long ago as 1608 by Blaise de Vigenere, in his Treatise on fire and salt, under the name of flowers of bensois, because it was obtained by sublimation; but it is now denominated benwoic acid.

1. The usual method of obtaining this acid is to put Preparaa quantity of benzoin, coarsely powdered, into an earthen pot, to cover the mouth of the pot with a cone of

thick paper, and then to apply a very moderate said heat. The benzoic acid is sublimed, and attaches itself This method was redious and difficult; to the paper. it being hardly possible to prevent the heat from scorching the benzoin, and volatilizing some empyreumie oil, which soils and injures the acid sublinied. Nezman proposed moistening the benzoin with alcohol, and distilling it in a retort with a low heat. comes over immediately after the alcohol, partly incrystals and partly of the consistence of butter. Ged. froy ascertained, in 1738, that this acid may be obtained by digesting benzoin in hot water. A portion is taken up, which is deposited in crystals as the water Scheele published a different method in 1775; which being easier and more productive than any of the preceding, is usually preferred. This process is as follows +: Upon four parts of unslacked lime pour twelve parts of water, and after the ebullition is over add 98 parts more of water; then put 12 parts of finely pounded benzoin into a tinned pan; pour upon it first about six parts of the above milk of lime, mix them well together, and thus successively add the rest of the mixture of lime and water. If it be poured in all at once, the benzoin, instead of mixing with it, will coagulate, and run together into a mass. This mixture ought to be boiled over a gentle fire for half an hour with constant agitation; then take it from the fire, let it stand quiet for an hour, in order that it may settle; pour off the supernatant limpid liquor into a glass vessel. Upon the remainder in the pan pour 96 parts of pure

^{*} Neuron's Chemistry, p. 294. † Scheele, i. \$24.

boil them together for half an hour, then take it from the fire, and let it settle; add the supernatent liquor to the former; pour upon the residuum some more water, boil it as aforesaid, and repeat the same process once more. At last put all the residuums upon a filter, and pour hot water several times upon it. During this process, the calcareous earth combines with the acid of benzoin, and separates it from the resinous particles of A small quantity of the resin is disthis substance. solved by the lime-water, whence it acquires a yellow All these clear yellow leys and decoctions are colour. to be mixed together, and boiled down to 24 parts; which are then to be strained into another glass vessel.

After they are grown cold, muriatic acid is to be added, with constant stirring, till there be no farther precipitation, or till the mass taste a little sourish. benzoic acid, which was before held in solution by the lime, precipitates in the form of a fine powder.

Mr Hatchett has observed, that when benzoin is digested in sulphuric acid, a great quantity of beautifully crystallized benzoic acid is sublimed. This process is the simplest of all, and yields the acid in a state of purity; it claims therefore the attention of manufacturers .

2. Benzoic acid, thus obtained, is a fine light whitish Properties. powder, which is not brittle, but has rather a kind of ductility. Its taste is acrid, hot, and somewhat bitter. Its odour is slight, but peculiar and aromatic †.

[#] Hatchett's Additional Experiments on Tannin. Phil. Trans. 1809.

[†] This odour is owing to a small portion of aromatic oil w ich adheres to the acid. It has been obtained without any smell by Giese, See Phil. Mag. ziv. 331.

specific gravity is 0.667 1. It hardly affects the infosion of violets, but it reddens that of turnsol, especially when hot ||.

Heat volatilizes this acid, and makes it give out strong odour, which excites coughing. When exposed to the heat of the blowpipe in a silver spoon, it melt, becomes as fluid as water, and evaporates without teking fire. It only burns when in contact with flame, and then it leaves no resideum behind. When thrown upon burning coals, it rises in a white smoke. When allowed to cool after being melted, it hardens, and a radiated crust forms on its surface ¶. When distilled in close vessels, the greater part of it sublimes unaltered, but some of it is decomposed. This portion is converted almost entirely into oil and carbureted hydroges

3. This acid is not altered by exposure to the air. Cold water dissolves no sensible quantity of it; but it is soluble enough in hot water: 480 grains of boiling water dissolves 20 grains of it; 19 of these are deposited, when the water cools in long, slender, flat, festher-like crystals ..

Action of simple bobies.

- 4. It is not affected by oxygen gas, not by any of the simple combustibles or incombustibles. It does not seem capable of oxidizing any of the metals +.
- 5. It combines with alkalies, earths, and metallic oxides, and forms salts known by the name of benzoates.
 - 6. Concentrated sulphuric acid dissolves it without

W Thid.

+ Trommederf.

T Lichte netein.

[!] Hassenfratz, Ann. de Chim. xxviii. 11

Morveau, Encyc. Method. Chim 1. 44.

beat, or any other change, except becoming somewhat brown: when water is poured into the solution, the benzoic acid separates, and coagulates on the surface without any alteration. Nitric acid presents precisely the same phenomena, as does also the sulphurous acid. Neither the muriatic, the oxymuriatic, nor the phosphoric acids dissolve it. Acetic acid, when hot, dissolves it precisely as water does; but it crystallizes again when the acid cools.

7. Alcohol dissolves it copiously, and lets it fall on the addition of water ‡. Boiling alcohol takes up its own weight §.

This acid is sometimes used as a medicine, but much less frequently than formerly.

SECT. III.

OF SEBACIC ACID.

CHEMISTS had long suspected that an acid could be History. obtained from tallow, on account of the acrid nature of the fumes which it emits at a high temperature; but it was M. Grutzmacher who first treated of it particularly, in a dissertation De Ossium Medulla, published in 1749 []. Mr Rhades mentioned it in 1753; Segner published a dissertation on it in 1754; and Crell exami-

^{*} Lichtenstein. † Id. † Id. § Wenzel's Verwandt-:chaft, p. 30%.

Leonhardi.

ned its properties very fully in two dissertations pub. lished in the Philosphical Transactions for 1780 and 1782. It was called at first acid of fat, and afterwards sebucic acid.

But at the period when these chemists made their experiments, the characteristic properties of the different acids were not sufficiently known to enable them to distinguish acids from each other with precision. Thenaid examined the subject in 1801, tried all the processes of Crell and Guyton Morveau, and found that the acids procured by them were either acetic or the acid employ. ed in the process. Real sebacic acid had hitherto escaped the examination of chemists. Thenard found, however, that a peculiar acid was formed during the distillation of tallow. To it he consigned the appellation of schacic acid. The experiments of this chemist were repeated in 1804 by Mr Rose, who obtained similar results, and confirmed all the observations of the French philosopher *. The subject has lately been resumed by Mr Berzelius, who in an elaborate dissertation, published in 1800, has shown that the sebacic acid of Thenard bears a close resemblance to benzoic acid+. Indeed he considers it as nothing else than benzoic acid contaminated with some unknown substance derived from the fat, which alters some of its properties, but from which it may be nearly freed by proper precautions.

Luchats-Çivil.

1. The method of procuring sebacic acid pointed out by Thenard is as follows: Distil hog's lard, wash the product with hot water, separate this water, and drop into

[#] Gehlen's Jour. iii, 170. | | 1 lbid. 2d Series, ii. 275.

it acctate of lead. A flaky precipitate appears, which is Chap. I be washed and dried, mixed with sulphuric acid, and heated. A melted substance, analogous to fat, swims on the surface, which is to be carefully separated. This substance is sebacic acid. It may be dissolved in hot water, and on cooling crystalline needles are deposited. This acid may be obtained also by evaporating the water employed in washing the product of distilled hog's lard. Or this water may be saturated with potash, and afterwards precipitated with acetate of lead as above .

During the distillation there comes over abundance of gas, which appears to be a mixture of carbonic acid and carbureted hydrogen. Into the receiver there drops a yellow-coloured water, impregnated with acetic acid, but containing no sebacic acid. It may, therefore, be put aside. But a much greater portion of oily matter passes over of the consistence of butter. The whole sebacic acid is contained in this matter, and may be separated by boiling it in water and evaporating the waacry solution. The sebacic acid falls down in the state of small crystals. From the experiments of Mr Rose, we learn that the quantity of sebacic acid formed by this process is extremely small; a pound of hog's lard yielding little more than 40 grains, and tallow or suct still less +.

2. Sebacic acid is white; it has no smell; its taste Proporties. is a pleasant sour, leaving in the mouth a very slight impression of bitterness. It reddens the infusion of turnsol. When heated it melts like tallow: on cooling it concretes into a crystalline mass. By a sudden and

^{*} Ann. de Chim. Exix. 194

Book IL Division IL violent heat it is decomposed. Berzelius has shown that it may be volatilized without decomposition, but that it requires a higher temperature for this purpose than benzoic acid.

Cold water dissolves but little sebacic acid, but boiling water takes up one-fourth of its weight. It separates as the water cools in small crystals. The crystals are needles; but by proper precautions it may be procured in long, large, and very brilliant plates. Alcohol dissolves it copiously, and so do the fixed and volatile oils. In all these respects, as Berzelius has shown, it agrees with benzoic acid.

It occasions a precipitate in the acetate and nitrate of lead, the nitrate of silver, the acetate and nitrate of mercury. It forms peculiar salts with the alkalies and It does not render lime-water, barytes, or strontian water, turbid. Sebate of potash has little taste, does not attract moisture from the air; and when sulphuric, nitric, or muriatic acid is poured upon it, sebacic acid is deposited: when its solution is concentrated and mixed with any one of these acids it becomes so-Berzelius has shown, that in these properties also sebacic and benzoic acid agree with each other. He has compared the different salts which each of these acids forms with the respective bases, and shown that they also nearly agree in their properties. Benzoic acid is usually combined with some extractive matter, from which it is nearly as difficult to free it as it is to free sebacic acid from the substances with which it is united. In short, from the experiments of Berzelius, there can

[#] Thenard and Rose, Ibid.

no doubt that the two acids, if not absolutely the Chap. II. ame, at least approach very nearly to each other.

SECT. IV.

OF SUCCINIC ACID.

AMBER is a well-known brown, transparent, inflam- History. mahir body, pretty hard, and susceptible of polish, found at some depth in the earth, and on the sea-coast of several countries. It was in high estimation among the ancients both as an ornament and a medicine.-When this substance is distilled, a volatile salt is obtained. which is mentioned by Agricola under the name of salt of amber; but its nature was long unknown. Boyle was the first who discovered that it was an acid *. From succinum, the Laun name of amber, this acid has received the appellation of succime acid.

Mr Pott seems to have been the first chemist who made a set of experiments to ascertain its properties, and who demonstrated that it differs from every other +.

1. It is obtained by the following process: Fill a re- Preparatort half-way with powdered amber, and cover the powder with a quantity of dry sand, lute on a receiver, and distil in a sand-bath without employing too much heat. There passes over first an insipid phlegm; then

Boyle abridged by Shaw, in. 369.

Men. relin. 1753 her an abstract of his paper in Lewis's edition of Neuman's Chemistry, p. 237.

a weak acid, which, according to Scheele, is the acetic then the succinic acid attaches itself to the neck of the retort; and if the distillation be continued, there comes over at last a thick brown oil, which has an acid taste.

The succinic acid is at first mixed with a quantity of oil. It may be made tolerably pure by dissolving it in hot water, and putting upon the filter a little conon, previously moistened with oil of amber: this substance retains most of the oil, and allows the solution to pass clear. The acid is then to be crystallized by a gentle evaporation; and this process is to be repeated till the acid be sufficiently pure. Guyton Morveau has shown that it may be made quite pure by distilling off it a sufficient quantity of nitric acid, taking care not to employ, a heat strong enough to sublime the succinic acid.

Properties.

2. The crystals of succinic acid are transparent, white, shining, and of a foliated, triangular, prismatic form: they have an acid taste, but are not corrosive: they redden tincture of turnsol, but have little effect on that of violets.

They sublime when exposed to a considerable heat, but not at the heat of a water-bath. In a sand-bath they melt, and then sublime and condense in the upper part of the vessel; but the coal which remains shows that they are partly decomposed ‡.

3. One part of this acid dissolves in 96 parts of water at the temperature of 50°, according to Spielman j; in 24 parts at the temperature of 52°; and in 2 parts of water at the temperature of 212°, according to Stock.

Bergman's Notes on Scheffer.

¹ Pott.

[†] Ann. de Chim. Exiz. E65.

[§] Inst. Chem. § zii,

de Neuforn*; but the greatest part crystallizes as water cools. According to Roux, however, it still tains more of the acid than cold water is capable of ssolving †.

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Two hundred and forty grains of boiling alcohol displue 177 of this acid; but crystals again shoot as the plution cools ‡.

- 4. Sulphuric acid dissolves it with the assistance of seat; but does not appear to decompose it. The same emark applies to nitric acid. Muriatic acid has but ittle action on it while cold; but when heat is applied, he whole coagulates into the consistence of a jelly \(\infty \).
- 5. The compounds which this acid forms with alkaies, acids, and metallic oxides, have received the name of succinates.
- Mhen combined with soda, it crystallizes in four and six sided prisms. When this salt is distilled in a retort, the succinic acid is completely decomposed. There passes over into the receiver an acid liquor, which is the sectic much diluted, and a quantity of brown oil. At the same time carbonic acid gas and carbureted hydrogen gas are disergaged, and their remain in the retort soda and charcoal ||. Hence it follows, that this acid, like the others of the same class, is decomposed by heat, and that it is composed of oxygen, hydrogen, and carbon.

De Succino. † Morveau, Encye. Method. Chim. i. 72.

[!] Wenzel's erwandtschaft, p. 305.
§ Pott.

Morveau, Ann. de Chim. xxix. 165.

SECT. V.

OF MOROXYLIC ACID.

History.

About the year 1802, Dr Thompson observed a saline exudation on the bark of the morus alba, or white mulberry tree, growing in the botanical garden of Palermo in Sicily. It coated the bark of the tree in small grains of a yellowish and blackish brown colour. A quantity of this matter was collected and sent to Mr Klaproth, who published an analysis of it in 1803.

It has a taste resembling that of succinic acid; and when placed upon burning coals, it emits a vapour which irritates the organs of smell. In water it dissolves speringly, forming a reddish-brown liquid, which yields by evaporation small needleform crystals, of the colour of pale wood, not deliquescing in the air. By dissolving these crystals in water, and treating them with various reagents, Klaproth ascertained them to be formed of lime united to a peculiar acid, which he called moroxylic, because the salt containing it was a production of the wood of the mulberry tree.

The salt dissolves but sparingly in water; hot water taking up only 0.035, and cold 0.015 of it. From the solution the alkaline carbonates precipitate lime; but bary tes causes no precipitate; a proof of the absence of

Scherer's Journal der Chemie, No. 55. p. 1. But I use the translation of the paper published in Nicholson's Journal, vii. 129.

silver, mercury, copper, iron, cobalt, and uranium, in tric acid; and of lead and iron in acetic acid. Baytes water, acetate of barytes, muriate of tin, muriate gold, and nitrate of nickel, produce scarcely any efect.

To obtain the acid, the solution of the salt was treatd with acetate of lead. Forty-five grains of the prepitate thus obtained were mixed with 20 grains of sulhuric acid diluted with 60 grains of water. The sulhate of lead formed remained in the state of a white
owder, while the moroxylic acid was dissolved and obnined by evaporation in the state of fine needles of the
colour of pale wood.

The acid thus obtained has the taste of succinic acid:
is not altered by exposure to the air: it dissolves really in water and in alcohol: it does not precipitate the etallic solutions like its salt.

When heated in a retort, a little acid liquor first times over, which has the taste of the concrete acid: text, the acid rises unaltered, and adheres to the top and eck of the retort in colourless and transparent prismatic crystals. A coaly residuum remains in the vessel. Hence it appears, that a gentle sublimation is the best mode of obtaining this acid in a state of purity.

When the calcareous salt of moroxylate of lime is istilled, a quantity of inflammable air mixed with carbonic acid is obtained, and an acid liquor comes over, which swims a brown oil. There remains in the setort a spongy mass of a pale brown colour, consisting of carbonate of lime mixed with some charry matter. From these phenomena it appears, that this acid, like the rest, is composed of oxygen, hydrogen, and carbon,

Chap. IL

Properties.

though the proportion of the constituents is unknown.

The compounds which it forms with alkalies have no ceived the name of moroxylates.

Such are the properties of this substance ascertained by Klaproth. The small quantity of the salt which he possessed prevented a more complete examination.

SECT. VI.

OF CAMPHORIC ACTO.

History.

Camphor is a well-known white crystalline substitics, of a strong taste and smell, obtained from a species of laurel in the East Indies. It is so volatile that it cannot be melted in open vessels, and so inflammable that it burns even on the surface of water.

When camphor is set on fire in contact with oxygen gas, it burns with a very brilliant flame; much caloric is disengaged, water is formed, the inner surface of the vessel is covered with a black matter, which is undoubtedly charcoal, and a quantity of carbonic acid gas is also produced *. Hence it follows, that it is composed of hydrogen and carbon, at least principally.

Mr Kosegarten, by distilling nitric acid off camphor eight times successively, obtained an acid in crystals to which the name of camphoric acid has been given.

Prepara-

1. His experiments have been repeated by Mr Bouil-

Bouillon La Grange, Ann. de Chim. xxiii. 153.

[†] Koecgarten & Campbera, &c. 1785.

La Grange. The process for obtaining camphoric d, as described by this chemist, is as follows: Put o a retort one part of camphor and eight parts of ni: acid, of the specific gravity 1.33. Distil with a d heat. A great deal of nitrous gas and carbonic 1 gas is emitted, and a little camphor sublimes. This cess must be repeated three times on the same campor; so that 24 parts in all of nitric acid are necessy. After the third distillation, when the retort is swed to cool, a number of crystals make their aparance in it. These are campboric acid. They aunt to somewhat less than the half of the camphor ployed.

2. Camphoric acid thus obtained is in snow-white Properties stals, of the form of parallelopipedons. These stals effloresce in the air †.

Camphoric acid has a slightly acid bitter taste, and a ell like that of saffron.

It reddens vegetable colours.

3. It is soluble in 200 parts of cold water, according Kosegarten; in 96 parts of water at the temperature 50°, according to Bouillon La Grange. Boiling wadissolves The thought The Transfer according to Bouillon La Grange.

According to Kosegarten, it is insoluble in alcohol; ording to Bouillon La Grange, alcohol dissolves it; I when the solution is left in contact with the air of atmosphere, the acid crystallizes. It is not preciated from its solution in alcohol by the addition of ter y.

Chap. H.

^{*} Kosegarten de Campbera, &c. 1785. † La Grange.

[!] Koscyarten. § Bouillon La Grange, Ann. de Chin. zivii. 40.

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Action of
heat.

4. When this acid is placed on ignited coals, it emits a dense aromatic fume, and is entirely dissipated. By a gentler heat it melts, and is sublimed. If it be put into a heated porcelain tube, and oxygen gas be passed through it, the acid does not undergo any change, but is sublimed.

By mere distillation it first melts, and then sublines; by which process its properties are in some respect changed. It no longer reddens the tincture of turnsol, but acquires a strong aromatic smell; its taste becomes less penetrating, and it is no longer soluble either in water or the sulphuric and muriatic acids. Heated nitric acid turns it yellow and dissolves it. Alcohol likewise dissolves it: and if this solution be left in contact with the air of the atmosphere, it crystallizes.

5. Camphoric acid does not produce any change on sulphur; alcohol and the mineral acids totally dissolve it; and so likewise do the volatile and the fat oils.

Camphoric acid does not precipitate lime from limewater. It produces no change on the solution of indigo in sulphuric acid.

6. It forms combinations with the alkalies, earths, and metallic oxides, which are called campborates.

SECT. VII.

OF OXALIC ACID.

When nitric acid is poured upon sugar, and a moderate heat applied, the sugar soon melts, an effervescence renes, a great quantity of autrous gas and carbonic soid is emitted; and when the effervescence ceases, and liquid in the retort is allowed to cool, a number of all transparent crystals appear in it. These crystals astitute a peculiar acid, which has received the name exalic acid, because it exists ready formed, as Scheele proved, in the oxalis acetosella, or wood-sorrel. At at, however, it was called the acid of sugar, or the echarine acid.

As the earliest and best account of the oxalic acid as published by Bergman, he was for a long time ckoned the discoverer of it; but Mr Ehrhart, one of cheele's intimate friends, informs us, that the world is debted for its knowledge of this acid to that illustris chemist *, and Hermbstadt and Westrumb assign the scovery to the same author +. The assertions of these ntlemen, who had the best opportunity of obtaining curate information, are certainly sufficient to establish e fact, that Scheele was the real discoverer of oxalic

1. Bergman gives us the following process for ob- Preparanoing this acid: " Put one ounce of white sugar powered into a tubulated retort, with three ounces of trong nitric acid, the specific gravity of which is to hat of water as 1.567. When the solution is over, duing which many fumes of the nitrous acid escape, let receiver be fitted, and the liquor made to boil, by which abundance of nitrous gas is expelled. When the liquor in the retort acquires a reddish-brown colour, dd three ounces more of nitric acid, and continue the

[·] Elwert's Magazine for Apothecories, 1785, part i. p. 54.

^{*} Kent's Distingary.

boiling till the fumes cease, and the colour of the lique vanishes. Then let the contents of the retort be emptied into a wide vessel; and, upon cooling, a crystallization will take place of slender quadrilateral prisms which are often affixed to each other at an angle of 45°. These crystals, collected and dried on blotting paper, will be found to weigh 12 dr. 19 gr. By boiling the remaining lixivium with two ounces of nitric acid in the retort till the red fumes almost disappear, and by repeating the crystallization as before, \$\dr. 13 gr. of solid acid will be obtained. If the process be repeated once more upon the residuum, which has now a glutinous consistence, with the successive additions of small quantities of nitric acid, amounting in all to two ounces, a saline brown deliquescent mass will be formed, weighing half a dram, of which about a half will be lost by a farther purification. The crystals obtained thus at different times may be purified by solution and crystallization, and by digesting the last lixivium with some nitric acid, and evaporation with the heat of the sun."

By the same process Bergman obtained it from gum arabic, alcohol, and honey. Scheele, Hermbstadt, Westrumb, Hoffman, &c. from a great variety of other vegetable productions; and Berthollet from a great number of animal substances.

It is of great consequence not to use too much nitric acid, otherwise the quantity of oxalic acid will be diminished; and if a very great quantity of nitric acid be used, no oxalic acid will be obtained at all. On the

[·] Bergman.

chap. IL mitric acid be used, the acid obtained will not be the scalic, but the tartaric. But I have not found this con-Fined upon trying the experiment. A 100 grains of sigar, when properly treated, yield 58 grains of crys-"Mized oxalic acid.

2. Oxalic acid, thus prepared, is in the form of four- Properties. med prisms, whose sides are alternately larger, and they are terminated at the extremities by two-sided summits. They are transparent, and of a fine white colour, with considerable lustre: they have a very acid taste, and redden vegetable blues. These crystals contain a portion of water, as happens to most crystallized bodies. When cautiously heated on a sand-bath they fall to powder, and lose about a third of their weight; but a part of this loss is to be ascribed to acid volatilied. From an accurate set of experiments made on purpose to investigate this point, I conclude them to be

23 water

77 real acid

100

somposed of

So that the water of crystallization amounts very nearly to the of the weight ".

3. When this crystallized acid is exposed to heat in Action of an open vessel, there arises a smoke from it, which affects disagreeably the nose and lungs. The residuum is a powder of a much whiter colour than the acid had been. By this process it loses 4d of its weight, but soon recovers it again on exposure to the air. When dis-

^{*} The reader is referred to a set of experiments by me on oxalic acid, published in the Phil. Trent. for 1807

tilled, it first loses its water of crystallization, then lique hes and becomes brown; a little phlegm passes over a white saline crust sublimes, some of which passed into the receiver; but the greatest part of the soid is destroyed, leaving in the retort a mass with of the whole, which has an empyreumatic smell, blackers sulphuric acid, renders nitric acid yellow, and dissolve in muriatic acid without alteration. That part of the acid which sublimes is unaltered. When this acid is distilled a second time, it gives out a white smole which, condensing in the receiver, produces a colourless uncrystallizable acid, and a dark coloured matter remains behind . During all this distillation a van quantity of clastic vapour makes its escape. From 279 grains of oxalic acid, Bergman obtained 109 cubic inches of gas, half of which was carbonic acid, and half carboreted hydrogen. Foutana from an ounce of it obtained 430 cubic inches of gas, one-third of which was carbonic acid, the rest carbureted hydrogen.

Of water.

4. The crystals of oxalic acid are soluble in their own weight of boiling water: water at the temperature of 65.7° dissolves half its weight of them. The specific gravity of the solution is 1.0593 †. One hundred parts of boiling alcohol dissolve 50 parts of these crystals; but at a mean temperature only 40 parts 1. Liquid oxalic acid has a very acrid taste when it is concentrated, but a very agreeable acid taste when sufficiently disluted with water §.

It charges all vegetable blues except indigo to a red. One grain of crystallized acid, dissolved in 1920 grains

@ Bergman. + Ibid, i. 255. 3 Ibal. 5 Ibid.

water, reddens the blue paper with which sugar eves are wrapt: one grain of it, dissolved in 3600 mains of water, reddens paper stained with turnsol *. According to Morveau, one part of the crystallized acid sufficient to communicate a sensible acidity to 2633 erts of water +.

Its fixity is such, that none of it is sublimed when exter containing it in solution is raised to the boiling perature.

5. Oxalic acid is not affected by exposure to the air, Of simple to the action of oxygen gas. The effect of the imple combustibles on it has not been tried; but in B probability is inconsiderable.

It is capable of oxidizing lead, copper, iron, tin, bisbuth, nickel, cobalt, zinc, manganese,

It does not act upon gold, silver, platinum, mercury.

- 6. Oxalic acid combines with alkalies, earths, and metallic exides, and forms salts known by the name of malates.
- 7. Muriatic and acetic acids dissolve oxalic acid, but without altering it 1. Sulphuric seid partly decompoit by the assistance of heat, and a quantity of charsoal is formed. Nitric acid decomposes it at a boiling beat, and converts it into water and carbonic acid f. From this result, and from the products obtained by distilling pure oxalic acid, it follows, that this acid is composed of oxygen, hydrogen, and carbon. The best method of ascertaining the proportion of these constituents, is to distil at a red heat determinate quantities

Chap. IL.

Bergman, i. 255.

Bergman.

[†] Bnaye. Method. art. Acide Sachaein.

⁵ Fourcroy, vii. 224.

of dry oxalutes. So far as my experiments have gon, the oxalic acid is decomposed precisely in the same way, and the same new products formed whatever onlate we use; but the oxalate of lime is most convenient, because we can ascertain its composition with the greatest accuracy. When exalate of lime is distilled in a small retort by means of a heat gradually raised to telness, the oxalic acid is completely decomposed and converted into five new substances; namely, water, carbone acid, carbonic oxide, carbureted bydrogen, and charcoil, The water is small in quantity, the carbonic acid large, and partly in the state of gas, partly combined with the base of the oxalate. The carbonic oxide and cubereted hydrogen are in the state of gases, and in the proportion of about 2; of the former to 1 of the latter. The charcoal is small in quantity, and mixed with the residue in the retort, to which it gives a grey colour. By carefully ascertaining the proportion of each of these products, and estimating their composition according to the most exact analysis hitherto made of each, I found that the composition of oxalic acid may be stated as follows, oxygen 64

Composi-

oxygen 64 carbon 32 hydrogen 4

100

This result differs very considerably from that announced by Fourcroy and Vauquelin as obtained by their experiments. Oxalic acid, according to them, is composed of oxygen 77, carbon 13, bydrogen 10. But I

Pourcroy, vii. 224.

persuaded that my numbers are much nearer the Chap. II. both than these. The proportion of carbon contained the carbonic acid, evolved by the distillation of oxate of lime, is a good deal greater than the whole of be carbon which they assign to the composition of oxae acid.

This acid is too expensive to be employed for the purposes of domestic economy; but it is extremely useal in chemistry to detect the presence of lime held in plution. For this purpose, either a little of the pure eid, or of the solution of oxalate of ammonia, is dropt sto the liquid supposed to contain lime. If any be resent, a white powder immediately precipitates. The reason of this is, that oxalate of lime is altogether psoluble, and oxalic acid in consequence is capable of aking lime from every other acid.

SECT. VIII.

OF MELLITIC ACID.

HERE is a mineral of a honey-yellow colour which History. found in small solitary crystals among the layers of Woodcoal at Arten in Thuringia. At first sight it has some resemblance to amber; but Werner recognised it

Provided the liquid does not contain an excess of any of the more powerful acids. In that case the excess must be saturated with ammois before adding the exalic acid.

Book II. Dividus II. as a peculiar substance about the year 1790, and gave it the name of bonigeteis (honey-stone), which foreign mineralogists converted into mellite. This mineral a very rare. Hitherto indeed it has been found only a Thuringia and in Switzerland.

Mellite has usually a honey-yellow colour, but sometimes a straw-yellow. It is always crystallized in octahedrons, but they are rarely entire; sometimes indeed almost the whole of one of the pyramids is wanting. Their surface is generally smooth and brilliant, and isteriorly they have a glassy lustre. They are semitrasparent, brittle, soft, and easily reduced to powder. When pounded, they assume a yellowish-grey colour. Their specific gravity is about 1.550 †.

Mineralogists soon discovered that mellite is partly combustible; but they did not agree about its component parts. Lampadius and Abich undertook its chemical analysis about the same time. According to the first of these gentlemen, it is composed of

86'4 charcoal

3.5 bitumen

2.0 silica

3.0 water

94.91

According to Abich, it contains the following ingre-

Brochant's Mineralogy, ii. 75.

⁺ Kispeoth's Bestrage, us. 245.

[†] Crell's Annals, 2797, 11. 20.

Chap. IL

40 carbonic acid

28 water

16 carbonate of alumina

54 benzoic acid

5 benzoate of alumina

3 oxide of iron

21 resin

100 +

The results of these analyses differed so much from each other, that little confidence could be placed in either. Besides, it was evident from the way in which their experiments were conducted, that the original component parts of mellite had been altered by fire. Klaproth analysed it in 1799, and ascertained it to be a compound of alumina and a peculiar acid, to which he gave the name of mellitic ‡. And this analysis was soon after confirmed by Mr Vauquelin ||.

1. Hitherto mellitic acid has been found only in the Preparamellite. It may be procured from that mineral by the following process: Reduce the mellite to powder, and boil it in about 72 times its weight of water. The acid combines with the water, and the alumina separates in flakes. By filtering the solution, and evaporating sufficiently, the mellitic acid is obtained in the state of crystals.

2. These crystals are either very fine needles, some- Properties. times collected into globules, or small short prisms.

[†] Crell's Annals, 1797, ii. p. 16.

[†] Beitrage, iii. 114.

Ann. de Chim. XXXVI. 203-

They have a brownish colour, and a taste at first sweet, ish-sour, and afterwards bitterish.

- 3. This acid is not very soluble in water; but the precise degree of solubility has not been ascertained.
- 4. When exposed to heat, it is readily decomposed, exhaling an abundant smoke, which however is destitute of smell. A small quantity of insipid ashes remain behind, which do not alter the colour of litmas paper.
- 5. All attempts to convert it into oxalic acid by the action of natric acid have failed. The nitric acid merely caused it to assume a straw-yellow colour.
- 6. The effect of the simple bodies on this acid has
- 7. It combines with alkalies, earths, and metallic orides, and forms with them salts which are distinguished by the name of mellates. The properties of them
 compounds will be considered afterwards.
- 6. From the analysis of Mr Klaproth, we learn that the mellite is composed of

46 mellitic acid

16 alumina

38 water

100

Now when mellite is distilled in a retort, the acid is completely decomposed, and its elements combine together in a different way. By this method Mr Klaproth obtained from 100 grains of mellite

54 cubic inches of carbonic acid gas

13 hydrogen gas

99 grains of acidulous and aromatic water

1 aromatic oil

. charcoal 16alumina

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From this analysis it is obvious, that mellitic acid is composed of carbon, hydrogen, and oxygen, like most of the combustible acids. It does not, however, furwish us with sufficient data for ascertaining the proportion of these constituent parts.

Mellitic acid bears a great resemblance to the oxalic; but it differs from it in many particulars, especially in the nature of the compounds which it forms with different bases.

SECT. IX.

OF TARTARIC ACID.

TARTAR, OF CREAM OF TARTAR, as it is commonly History. called when pure, has occupied the attention of chemists for several centuries. Duhamel and Grosse, and after them Margraff and Rouelle the Younger, proved that it was composed of an acid united to potash; but Scheele was the first who obtained this acid in a separate state. He communicated his process for obtaining it to Retzius, who published it in the Stockholm Transactions for 1770. It consisted in boiling tartar with chalk, and in decomposing the tartrate of lime thus formed by means of sulphutic acid.

1. The process employed at present for obtaining tar- Preparataric acid, which is the same with that of Scheele, is the following: Dissolve tartar in boiling water, and add

to the solution powdered chalk till all effervescence ceases, and the liquid does not redden vegetable bluss Let the liquid cool, and then pass it through a filter. A quantity of tartrate of lime (which is an insoluble white powder) remains upon the filter. Put this tar. trate, previously well washed, into a glass cucurbite, and pour on it a quantity of sulphuric seld equal to the weight of the chalk employed, which must be diluted with water. Allow it to digest for 12 hours, stirring it occasionally. The sulphuric seid displaces the tartarte: sulphate of lime remains at the bottom, while the tartaric acid is dissolved in the liquid part. Decant off this last, and try whether it contains any sulphurie acid. This is done by dropping in a little acetate of lead; a precipitate appears, which is insoluble in acetic acid if sulphuric acid be present, but soluble if it be ab-If sulphuric acid be present, the liquid must be digested again on some more tartrate of lime; if not, it is to be slowly evaporated, and about one third part of the weight of the tartar employed is obtained of crystallized tartaric soid.

Lime may be substituted for chalk in this process. In that case the decomposition of the tartar is complete; whereas by Scheele's method, the excess of acid only combines with the chalk; but when lime is used, the whole tartrate of lime by no means separates. A considerable portion is retained in solution by the potash of the tartar now disengaged. If the liquid be evaporated, this portion appears under the form of a transparent jelly. By exposure to the air the potash attracts carbonic acid, which unites to the lime, while the tartaric acid combines again with the potash. To obtain the potash in a state of tolerable purity, the best me-

ad, according to Vauquelin, to whom we are indebtfor these observations, is to evaporate to dryness, and the residue to redness. By lixiviating the mass, s potash will be obtained in a state of considerable

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2. The form of the crystals of tartaric acid is so ir- Properties. gular, that every chemist who has treated of this subet has given a different description of them. Accordg to Bergman, they generally consist of divaricating mellat +; according to Van Packen, they assume oftz est the form of long-pointed prisms 1; Spielman and Corvinus i obtained them in groups, some of them nce-shaped, others needle-formed, others pyramidal. Morveau obtained them needle-form | . Their specific avity is 1.5962 ¶.

As they are sold by apothecaries in this country, they e in groups, and appear to be very irregular four-sided By my experiments they are composed of 14.5 real acid and 15.5 of water.

3. Crystallized tartaric acid is white, and very firm, is but imperfectly transparent. It may be exposed the air for any length of time without undergoing any lange. When heated a few degrees above 2120 it celts and remains limpid and transparent like water. 1250° it boils without losing its transparency or beming coloured. If the boiling be continued for a wort time only it does not lose above 4 per cent. of its

[·] Ann. de Chim, zlvil. 147.

[†] Bergman, iii 368.

¹ De Sale Frent. Acidi Tortari.

Analesta de Turare.

Eneye. Method. Chim. i. 323.

Hamenfrats, Ann. de Chim, xxxviii. It

Book 17. Division 11. weight. On cooling it concretes again into a hard we mitransparent mass, very much resembling a quantity of white sugar that has been melted at the same temperature. By this process the nature of the acid is changed. It now deliquences when exposed to the air.

In the open fire tartaric acid burns without leaving any other residuum than a spongy charcoal, which generally contains a little lime. When distilled in close vessels, it is converted into carbonic acid gas and carbonic acid hydrogen gas, a coloured oil, and a reddish scilliquor, which was formerly distinguished by the name of pyrotartarous acid, but which Fourcroy and Vauquelin have lately ascertained to be merely acetic acid impregnated with oil.

When tartaric acid combined with a base, as with lime, is distilled in a sufficiently strong heat, it is completely decomposed. The products into which it is converted, are, water, carbonic acid, beavy inflammable air, oil and charcoal. The quantity of carbonic acid which it yields is less than what is furnished by an equal weight of oxalic acid; but it differs from oxalic acid, in yielding a portion of brown coloured thick oil, which has an empyreumatic smell, and dissolves in alcohol.

4. Tartaric acid dissolves readily in water. Bergman obtained a solution, the specific gravity of weich was 1.230†. Morveau bserved, however, that crystals formed spontaneously in a solution, the specific gravity of which was 1.084. It is not liable to spec-

Bergman, i. 250.

[&]quot; Ann. de Chim. 2424, 141.

meous decomposition when dissolved in water, unless Chap. It. solution be considerably diluted.

5. Neither its action on oxygen gas nor on simple embustibles and incombustibles has been examined; Action of at it is probable that it is not capable of producing any dies. nsible change on them. It is capable of oxidizing iron d zinc, and even mercury; but it does not act upon timony, bismuth, tin, lead, copper, silver, gold, nor latinum. Its action on the other metallic bodies has procly been examined.

6. It combines with alkalies, earths, and metallic oxes, and forms salts known by the name of tartrates. Though it forms with the alkaline earths salts which e scarcely soluble in water, yet it has not the properof precipitating any of them from their solutions. this respect it differs very greatly from oxalic mid, which precipitates them all except magnesia.

7. The action of the greater part of the other acids a it is unknown. Hermbstadt has ascertained, that it my be converted into oxalic acid by distilling it repeatdly with six times its weight of nitric acid. By this process he obtained 560 parts of oxalic acid from 360 erts of tartaric acid 1.

8. From this result, and from the products obtained then tartaric acid is distilled, it is evident that it is Composicomposed of oxygen, carbon, and hydrogen. Fourcroy oforms us, that Vauquelin and he have ascertained but these ingredients are combined in it in the followproportions :

70·5 oxygen 19·0 carbon 10·5 hydrogen

100.0

Put to any use; but some of the compounds into which it enters are much employed in medicine. This said has the property of combining in two different propositions with a great number of bases. With potash, for instance, in one proportion, it forms a sait pretty soluble in water, called tartrate of potash; but when added in a greater proportion, it forms tartar, a sait very many perfectly soluble in water. By this property, the presence of tartaric acid in any acid solution may can't be detected. All that is necessary is to drop in slowly a little solution of potash: if tartaric acid be present tartar immediately precipitates in the form of a white gritty powder.

SECT. X.

OF CITRIC ACID.

History.

CHEMISTS have always considered the juice of oranges and lemons as an acid. This juice contains a quantity of mucilage and water, which renders the acid in pure, and subject to spontaneous decomposition. Mr Georgius took the following method to separate iciolage. He filled a bottle entirely with lemon-juice, cork-

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ed it, and placed it in a cellar: in four years the liquid was become as limpid as water, a quantity of mucilage had failen to the bottom in the form of flakes, and a thick crust had formed under the cork. He exposed this acid to a cold of 23°, which froze a great part of the water, and left behind a strong and pretty pure acid . It was Scheele, however, that first pointed out method of obtaining this acid perfectly pure, and who demonstrated that it possesses peculiar properties.

1. His process, which is still followed, is this: Satu- Prepararate lemon juice, while boiling, with powdered chalk. A white powder falls to the bottom, which is lime combined with citric acid. Separate this powder, and wash it with warm water till the water passes off cotourless. Then pour upon it as much sulphuric acid as will saturate the chalk employed, having previously diluted it with six times its weight of water. Boil the mixture for some minutes, and pass it through a filter to separate the sulphace of lime. The liquid is then to be evaporated to the consistence of a syrup, and set aside to cool. A number of crystals form in it: these use citric acid t.

Mr Scheele advises the use of an excess of sulphuric acid in order to ensure the separation of all the lime; but according to Dize, this excess is necessary for another purpose 1. A quantity of mucilage still adheres to the cirric acid in its combination with lime, and sulphuric acid is necessary to decompose this mucilage a which, as Fourcroy and Vauquelin have proved, it is

Stockbolm Tres ations, 1774. 1 N cholson's Journal, ii, 43.

† Scheele, il. 203.

capable of doing. His proof of the presence of much lage is, that when the solution of citric acid in water, which he had obtained, was sufficiently concentrated by evaporation, it assumed a brown colour, and even became black towards the end of the evaporation. The crystals also were black. By repeated solutions and svaporations, this black matter was separated, and found to be charcoal. Hence he concluded that mucilage had been present; for mucilage is composed of carbon, hydrogen, and oxygen; sulphuric acid causes the hydrogen and oxygen to combine and form water, and chart coal remains behind. It is not certain, however, as Me Nicholson remarks very justly *, that the sulphuric scil may not act upon the citric acid itself, and that the charcoal may not proceed from the decomposition of it; at least the experiments of Mr Dizé are insufficient to prove the contrary. In that case, the smaller the excess of sulphuric acid used the better.

Proust, who has published a memoir on the preparation of citric acid in the Journal de Physique for 1801; has observed that this is actually the case. When too much sulphuric acid is employed, it acts upon the citric acid, chars it, and prevents it from crystallizing. This error is remedied by adding a little chalk. This chemist has ascertained, that four parts of chalk require for saturation 94 parts of lemon juice. The citrate of lime obtained amounts to 7; parts. To decompose this, 20 parts of sulphuric acid, of the specific gravity 1.15, are necessary.

Properties.

2. The crystals of citric acid are rhomboidal prisms,

[·] Nicholson's Journal, ii. 43.

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about 120 and 60 degrees, terminated at each end by ar trapezoidal faces, which include the solid angles. hey are not altered by exposure to the air. Their use is exceedingly acid, and even excites pain; but hen the acid is properly diluted with water, the acidity pleasant.

- 3. When exposed to the open fire, it first melts, then rells up and exhales an acrid vapour, and leaves beand it a small quantity of charcoal. When distilled in ose vessels, it partly evaporates without decomposition, d is partly converted into acetic acid, carbonic acid, d carbureted hydrogen gas, which pass over, and arcoal which remains in the retort.
- 4. Citric acid is exceedingly soluble in water. According to Vauquelin, 100 parts of it require only 75 texts of water. Boiling water dissolves twice its weight it †. This solution may be kept a long time in close tessels; however it at last putrifies, and is decomposed.
- 5. Neither oxygen gas nor the simple combustibles or incombustibles produce any effect upon it. It is capable of oxidizing iron, zinc, tin. It does not act upon cold, silver, platinum, mercury, bismuth, antimony, arrenic.
- o. It combines with alkalies, earths, and metallic oxides, and forms salts known by the name of citrates.
- 7. The action of none of the acids on it has been exunined, if we except that of the sulphuric and nitric. Sulphuric acid, when concentrated, converts it into a-

Action of

Scheele said that he could not convert it cetic acid *. into oxalic acid by means of nitric acid, as he had done several other acids: but Westrumb affirms, that this conversion may be affected; and thinks that Scheele had probably failed from having used too large a quantity of nitric acid, by which he had proceeded beyond the conversion into oxalic acid, and had changed the citric acid into vinegar: and in support of his opinion, he quotes his own experiments; from which it appeared that, by treating 60 grains of citric acid with different quantities of nitric acid, his products were very different. Thus with 200 grains of nitric acid he got 30 grains of oxalic acid; with 300 grains of nitric acid he obtained only 15 grains of the oxalic acid; and with 600 grains of nitric acid no vestige appeared of the oxalic On distilling the products of these experiments, especially of the last, he obtained vinegar mixed with nitric acid. The experiments of Westrumb have been confirmed by Fourcroy and Vauquelin; who, by treating citric acid with a great quantity of nitric acid, converted it into oxalic and acetic acids. The proportion of the first was much smaller than that of the second.

From these experiments it is evident that its component parts are oxygen, hydrogen, and carbon; but the proportions of them have not been ascertained.

The use of this acid as a seasoner of food, and in preparing an agreeable cooling drink called *lemonade*, is well known.

Fourcroy, vii. 206.

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SECT. XI.

OF KINIC ACID.

MR DESCHAMPS, junior, an apothecary in Lyons, some Discovery. time ago published a method of extracting from yellow Peruvian bark a peculiar salt, to which the physicians of Lyons ascribed the febrifuge properties of that bark. His process was very simple; the bark was macerated in cold water, and the infusion concentrated by evapo-It was then set aside for some time in an open The crystals of the salt gradually formed and separated, and they were purified by repeated crystallizations. From 100 parts of the bark about 7 parts of these crystals were obtained *.

This salt has been lately examined by Mr Vauquelin, who has proved that it is composed of lime united to a peculiar acid hitherto unknown. To this acid he has given the name of kinic, borrowed from the term quinquina, applied by the French writers to the bark from which the salt was extracted.

Deschamp's salt then is kinate of lime. It possesses Kinate of the following properties.

lime.

Its colour is white, it crystallizes in square or rhomboidal plates; it has no taste, and is flexible under the teeth. It dissolves in about five times its weight of

^{*} Ann. de Chim. xlviii. 162.

water at the temperature of 55°. It is insoluble a alcohol.

When placed on burning coals it swells, gives out a smell like tartar, and leaves a mixture of carbonate of lime and charcoal. The fixed alkalies and their carbonates precipitate lime from the solution of this sale, but ammonia produces no effect. Sulphuric and oxalic acid precipitate lime likewise, but no precipitate is produced by acetate of lead or nitrate of silver. The infusion of tannin occasions a flaky yellow precipitate.

Separation of kinic soid. Mr Vauquelin separated the lime from the kinic acid by means of oxalic acid. The oxalate of lime obtained from 100 grains of the salt weighed 27 grains. Hence he concluded that the salt was a compound of \$5 parts acid and 15 lime. But the lime in 27 grains of oxalate of lime very little exceeds 10 grains. Hence the salt ought to be composed of 20 parts acid and 18 of lime.

Properties.

The kinic acid, thus freed from lime, was concentrated by evaporation to the consistency of a syrup, and then set aside for a week. No crystals formed in it spontaneously; but upon being touched with a glast rod it wholly crystallized at once in divergent plates.

Its colour was slightly brown (doubtless from some impurity); its taste was extremely acid and somewhat bitter, probably because it was not quite free from the other constituents of the bark. It was not altered by being left exposed to the air.

On burning coals it melted, froathed, blackened, and exhaled in acrid vapours, leaving only a little charcoal behind.

It combines with the different bases, and forms a set of salts called kinates. The alkaline and earthy ki-

mates are soluble and crystallizable. This ac does Chap. II.

This detail, though imperfect, is sufficient to disinguish the kinic from every other acid. Its decomposition by heat shows that it belongs to the class of combastible acids. Its crystallizing, its great solubility in water, and its forming a soluble salt with lime, and not precipitating silver nor lead, sufficiently distinguish is from all the other combastible acids with which it might be confounded.

SECT. XII.

OF SACLACTIC ACID.

This acid was discovered by Scheele in 1780. After having obtained oxalic acid from sugar, he wished to examine whether the sugar of milk would furnish the same product. Upon four ounces of pure sugar of milk, finely powdered, he poured 12 ounces of diluted nitric acid, and put the mixture in a large glass retort, which he placed in a sand-bath. A violent effervesence ensuing, he was obliged to remove the retort from the sand-bath till the commotion ceased. He then continued the distillation till the mixture became yellow. As no crystals appeared in the liquor remaining in the retort after standing two days, he repeated the distilla-

[·] Ann. de Chim. lix. 162.

tion as before, with the addition of eight ounces of mitric acid, and continued the operation till the yellow colour, which had disappeared on the addition of the aitric acid, returned. The liquor in the retort contamed a white powder, and when cold was observed to be thick. Eight ounces of water were added to dilute this liquor, which was then filtrated, by which the white powder was separated; which being edulcorated and dried, weighed 7 dr. The filtrated solution was evaporated to the consistence of a syrup, and again sub. jected to distillation, with four ounces of nitric acid as before; after which, the liquor, when cold, was observed to contain many small, oblong, sour crystals, together with some white powder. This powder being separated, the liquor was again distilled with more mitric acid as before; by which means the liquor was rendered capable of yielding crystals again; and by one distillation more, with more nitric acid, the whole of the liquor was converted into crystals. These crystals. added together, weighed five drams; and were found, upon trial, to have the properties of the oxalic acid.

Mr Scheele next examined the properties of the white powder, and found it to be an acid of a peculiar nature; he therefore called it the acid of the sugar of milk. It was afterwards called saclactic acid by the French chemists. Fourcroy has lately given it the name of mucous acid, because it is obtained by treating gum arabic, and other mucilaginous substances, with nitrigacid.

[♥] Setteele, ii, 69.

Chap. It.

Mr Hermbstadt of Berlin had made similar experiments on sugar of milk at the same time with Scheele, and with similar results; but he concluded that the white powder which he obtained was nothing else than exalate of lime with excess of acid, as indeed Scheele himself did at first. After he became acquainted with Scheele's conclusions, he published a paper in defence of his own opinion; but his proofs are very far from establishing it, or even rendering its truth probable ". He acknowledges himself, that he has not been able to decompose this supposed salt; he allows that it possesses properties distinct from the oxalic acid; but he ascribes this difference to the lime which it contains : yet all the lime which he could discover in 240 grains of this salt was only 20 grains; and if the alkali which he employed was a carbonate (as it probably was), these 20 must be reduced to 11. Now Morveau has shown, that oxalic acid, containing the same quantity of lime, exhibits very different properties. Besides, this acid, whatever it is, when united with lime, is separated by the oxalic, and must therefore be different from it. as it would be absurd to suppose that an acid could displace itself +. The saclactic acid must therefore be considered as a distinct acid, since it possesses peculiar properties.

1. Saclactic acid may be obtained by the following Prepuraprocess: Upon one part of gum arabic, or other similar gum, previously put into a retort, pour two parts of nitric acid. Apply a slight heat for a short time, till a

Crell'e Annals, 1784, il. 509.

Morvent, Encyc. Metbod, i. 291.

little nitrous gas and carbonic acid gas comes over; then allow the mixture to cool. A white powder gradually precipitates, which may be easily separated by filtration. This powder is saclactic acid.

Properties.

2. Saclactic acid, thus obtained, is under the form of a white gritty powder, with a slightly acid taste.

Heat decomposes it. When distilled, there comes over an acid liquor which crystallizes in needless on cooling, a red coloured acrid oil, carbonic acid gas, and carboneted hydrogen gas. There remains in the retort a large proportion of charcoal.

Saclactic acid, according to Scheele, is soluble in 60 parts of its weight of boiling water; but Messrs Hermbstadt † and ‡ Morveau found, that boiling water only dissolved to the part: it deposited about 4th part on cooling in the form of crystals j.

The solution has an acid taste, and reddens the infusion of turnsol ||. Its specific gravity, at the temperature of 53.7°, is 1.0015 ¶.

The compounds which it forms with earths, alkalies, and metallic oxides, are denominated saccolates. These salts are but very imperfectly known, no chemist having examined them except Scheele. The facts which he ascertained are the following:

With potash it forms a salt soluble in eight parts of boiling water, which crystallizes on cooling. The saccolate of soda also crystallizes, and is soluble in five parts of water. Saccolate of ammonia loses its base by

۴ Id.

† Pipe. Co

^{*} Fourcroy, vii. 146. ‡ Encyc. Method. i. 290.

cate heat, while the acid remains behind. The Chap. It. ands which it forms with the earths are nearly he in water. It scarcely acts upon any of the but it combines with their oxides: it forms most insoluble in water. When poured into the s of silver, mercury, or lead, it occasions a white tate. It produces no change in the sulphates of opper, zinc, manganese, nor in the muriates of mercury ".

SECT. XIII.

OF URIC ACID.

ous attempts were made by chemists to ascertain Discovery. cure of the calculous concretions which occasionin the kidneys and bladder, and produce one most painful diseases to which we are liable. attempts were attended with very little success, beele published a set of experiments on the sub-1776. He examined several of these urinary and found them composed chiefly of a peculiar be properties of which he described. His resul boon after confirmed by Bergman, who had ena similar set of experiments about the same To the acid thus discovered, Morveau gave a name of bezoardic, which was after changed i

1 Ibid. i. 299 and 210. French Trans

to lithic by the French chemists, when they contrived the new chemical nomenclature in 1787. This last term, in consequence chiefly of the observations and objections of Dr Pearson, has been recently laid aside, and the name uric acid substituted in its place.

Scheele ascertained that uric acid exists always in human urine. Experiments on the urinary calculi were published by Mr Higgins in 1789 *, and by Dr Austin in 1791; but little was added to our knowledge of uric acid till Dr Wollaston published his admirable paper on the calculous concretions in 1797. Dr Pearson published a copious set of experiments on the same subject in 1798, in which he enumerates his trials on uric acid, and endeavours to prove that it is not entitled to the name of acid, but ought to be classed among animal oxides. This drew the attention of Fourcroy and Vauquelin to the subject: they published an excellent treatise on urinary calculi, in which they demosstrate that it possesses the properties of an acid, and confirm the observations of Bergman and Scheele. But for the most complete account of uric acid we are endebted to Dr William Henry, who made it the subject of his thesis published in 1807.

Prepara-

1. To obtain pure uric acid, Dr Henry dissolved pulverized calculi (previously known to be composed chiefly of that acid) in a ley of potash, and precipitated the uric acid by means of muriatic or acetic acids. The powder thus obtained was first washed with a little ammonia, to remove any adhering foreign acid, and

Comparative view of the phlogistic and antiphlogistic theories, p. 42.

en edulcorated with a sufficient quantity of warm Chap II.

2. Uric acid thus obtained is a white powder, which properties harsh but not gritty, and is destitute both of taste and smell. It reddens the infusion of litmus. It dissives in 1720 parts of water at the temperature of 60°, and in 1150 parts of boiling water. As this last solution cools, it deposites minute crystals of uric acid. The watery solution reddens litmus, but produces no tecrpitate when dropt into earthy or metallic salts.

S. It is rapidly dissolved by fixed alkaline solutions, at less readily by ammonia. The alkaline carbonates are no effect on it whatever †.

4. It decomposes the alkaline hydrosulphurets, and precipitates their sulphur. It likewise decomposes on provided a sufficient quantity of it be employed 1.

5. It dissolves in nitric acid; and when the solution is evaporated nearly to dryness, it assumes a fine pink colour, which becomes much deeper when water is added, so as to have a near resemblance to carmine. In this state it stains wood, the skin, &c. of a beautiful recolour. The watery solution of this matter loses its red colour in a few hours, and it cannot afterwards be restored §.

When the solution of uric in nitric acid is boiled, a quantity of azotic gas, carbonic acid gas, and of prustic acid, is disengaged ||. Dr Pearson, by repeated distillations, converted the residue into nitrate of ammonia. When oxymuriatic acid gas is made to pass into water

Henry. † Scheele,

[†] Scheele, fourcroy, and Henry.

1 Behaline nan, Pearson, and Henry.

t Heavy.

containing this acid suspended in it, the acid assumes a gelatinous appearance, then dissolves; carbonic scid gas is emitted, and the solution yields by evaporation muriate of ammonia, superoxalate of ammonia, muriate acid, and malic acid.

- 6. It combines with the different bases, and forms a genus of salts called urates; for the examination of which we are chiefly indebted to Dr Henry.
- 7. When uric acid is distilled, about a fourth of the acid passes over altered in its properties, and is found in the receiver crystallized in plates; a few drops of thick oil make their appearance; ith of the acid of concrete carbonate of ammonia, some prussiate of ammonia, some water, and carbonic acid pass over; and there remains in the retort charcoal, amounting to about ith of the weight of the acid distilled. According to De Henry, the decomposition goes on in the following manner: 1. A drop or two of water holding carbonate of ammonia in solution passes into the receiver; 2. Dry carbonate of ammonia; 3. The acid sublimate; 4. The coal in the retort amounts to about the calculus distilled.

Sublimate from uric acid. The nature of the acid sublimate has been particularly examined by Dr William Henry. Scheele had considered it as analogous to succinic acid, and Dr Pearson as approaching to benzoic acid in its properties.
According to Dr Henry, it is a salt composed of a proculiar acid combined with ammonia. The following
he found to be its properties.

^{*} Brugnstelli, Ann. de Com. 22vin 267 Four roy, 2. 244.

⁺ Fourcroy, dan. de Chim ave 216.

Its colour is yellow, and it has a cooling bitter taste. It dissolves readily in water, and in alkaline solutions, from which it is not precipitated by acids. It dissolves aso sparingly in alcohol. It is volatile; and when ablimed a second time, becomes much whiter. The watery solution reddens vegetable blues; but a very mall addition of ammonia destroys this property. It des not produce an effervescence with alkaline carbon-By evaporation it yields permanent crystals, but defined, from a portion of animal matter adhering. They redden vegetable blues. Potash, when added to these crystals, disengages ammonia. When dissolved nitric soid they do not leave a red stain, as happens with uric acid; nor does their solution in water decompose the earthy salts, as happens with the alkaline grates: neither has it any action on the salts of copper, fron, gold, platinum, tin, or mercury. With nitrates of silver and mercury, and acetate of lead, it forms a white precipitate soluble in an excess of nitric acid, Muriatic acid occasions no precipitate in the solution of these crystals in water. These properties show us, that the acid of the sublimate is different from the uric. and from every other known acid. Dr Austin found, that by repeated distillations it was resolved into ammonia, azote, and prussic acid. Hence its constituents are doubtless the same as those of uric acid, varying only in proportion.

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SECT. XIV.

OF LACCIC ACID.

Mistory.

About the year 1786, Dr Anderson of Madras mentioned, in a letter to the governor and council of that place, that nests of insects, resembling small cowry shells, had been brought to him from the woods by the natives, who eat them with avidity. These supposed nests he soon afterwards discovered to be the coverings of the females of an undescribed species of coccus, which he shortly found means to propagate with great facility on several of the trees and shrubs growing in his neighbourhood.

On examining this substance, which he called white lac, he observed in it a very considerable resemblance to bees wax; he noticed also, that the animal which secretes it provides itself by some means or other with a small quantity of honey, resembling that produced by our bees; and in one of his letters he complains, that the children whom he employed to gather it were tempted by its sweetness to eat so much of it as materially to reduce the product of his crop. Small quantities of this matter were sent into Europe in 1789, both in its

The Chinese collect a kind of wax, which they call pe-la, from a coccus, deposited for the jurpose of breeding on a viral shrubs, and manage it exactly as the Mexicans manage the co-hineal usect. It was the knowledge of this that induced Dr Anderson to attempt to propagate his insect.

Extural state and melted into cakes; and in 1793 Dr Extraon, at the request of Sir Joseph Banks, undertook Chemical examination of its qualities, and his experitents were published in the Philosophical Transactions 1794. Chap. II.

A piece of white lac, from 3 to 15 grains in weight, probably produced by each insect. These pieces are f a grey colour, opaque, rough, and roundish. When thite lac was purified by being strained through musin, it was of a brown colour, brittle, hard, and had a itterish taste. It melted in alcohol, and in water of the temperature of 145°. In many of its properties it exembles bees wax, though it differs in others; and Ir Pearson supposes that both substances are composed f the same ingredients, but in different proportions.

1. Two thousand grains of white lac were exposed in Properties, uch a degree of heat as was just sufficient to melt nem. As they grew soft and fluid, there oozed out 50 grains of a reddish watery liquid, which smelled ke newly baken bread *. To this liquid Dr Pearson as given the name of of laccic acid †.

2. It possesses the following properties:

It turns paper stained with turnsol to a red colour.

After being filtered, it has a slightly saltish taste with itterness, but is not at all sour.

When heated, it smells precisely like newly baken hot read.

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^{*} The same liquid appears on pressing the crude lac between the ngers; and we are told that when newly gathered it is replete with ice.

[†] Pearson's Transl. of the Clemical Nomenclature.

On standing, it grows somewhat turbid, and deposits a small quantity of sediment.

Its specific gravity at the temperature of 60° is 1.025.

A little of it having being evaporated till it grew very turbid, afforded on standing small needle-shaped crystals in mucilaginous matter.

Two hundred and fifty grains of it were poured into a very small retort and distilled. As the liquor grew warm, mucilage-like clouds appeared; but as the head increased they disappeared again. At the temperature of 200° the liquor distilled over very fast; a small quantity of extractive matter remained behind. The distilled liquor while hot smelled like newly baken bread, and was perfectly transparent and yellowish. A shred of paper stained with turnsol, which had been put into the receiver, was not reddened; nor did another which had been immersed in a solution of sulphate of iron, and also placed in the receiver, turn to a blue colour upon being moistened with the solution of potash.

About 100 grains of this distilled liquid being evaporated till it grew turbid, after being set by for a night, afforded acicular crystals, which under a lens appeared in a group not unlike the umbel of parsley. The whole of them did not amount to the quarter of a grain. They tasted only bitterish.

Another 100 grains being evaporated to dryness in a very low temperature, a blackish matter was left behind, which did not entirely disappear on heating the spoon containing it very hot in the naked fire; but on

[•] A proof that the acid was not the prussic.

oxalic acid to a much less degree, it evaporated, Chap. II., t not a trace behind.

onate of lime dissolved in this distilled liquid with The solution tasted bitterish, did not aper stained with turnsol red, and on adding to onate of potash a copious precipitation ensued. A f this solution of line and of alkali being eval to dryness, and the residuum made red hot, noemained but carbonate of lime and carbonate of

i liquid did not render nitrate of lime turbid, but aced turbidness in nitrate and muriate of barytes. 500 grains of the reddish-coloured liquor obtainnelting white lac, carbonate of soda was added till rvescence ceased, and the mixture was neutrafor which purpose three grains of the carbonate ecessary. During this combination a quantity ilaginous matter, with a little carbonate of lime, ecipitated. The saturated solution being filtral evaporated to the due degree, afforded on standliquescent crystals, which on exposure to fire left residuum of carbonate of soda.

e-water being added to this teddish-coloured liproduced a light purple turbid appearance; and on g there were clouds just perceptible.

huret of lime occasioned a white precipitation, sulphureted hydrogen gas was perceptible by the

ture of galls produced a green precipitation. hate of iron produced a purplish colour, but no tation; nor was any precipitate formed by the n first of a little vinegar, and then of a little potthe mixture.

Acetate of lead occasioned a reddish precipitation, which redissolved on adding a little natric acid.

Nitrate of mercury produced a whitish turbid li-

Oxalic acid produced immediately the precipitation of white acicular crystals, owing probably to the presence of a little lime in the liquid.

Tartrate of potash produced a precipitation not unlike what takes place on adding tartaric acid to tartrate of potash; but it did not dissolve again on adding potash.

Such were the properties of this acid discovered by Dr Pearson*. Before this acid can be admitted as a peculiar substance, it would be necessary to subject it to a more rigid ex mination. The quantity which Dr Pearson examined was by far too small to enable him to obtain satisfactory results.

SECT. XV.

OF MALIC ACID.

Preparation. 1. This acid, which was discovered by Scheele in 178, has received the name of malic acid, because it may be obtained in abu dance from the juice of apples, in which it exists ready formed. Scheele has given us the following process for extracting it: Saturate the juice of apples with potash, and add to the solution ace-

^{*} Pbil. Tress. 1794, p. 383.

the of lead till no more precipitation ensues. Wash he precipitate carefully with a sufficient quantity of vater; then pour upon it diluted sulphuric acid till the nixture has a perfectly acid taste, without any of that weetness which is perceptible as long as any lead renains dissolved in it; then separate the sulphate of ead, which has precipitated, by filtration, and there renains behind pure malic acid.

Vauquelin has lately ascertained that it may be exracted, with greater advantage, from the juice of the tempervivum tectorum, or common house-leek, where it exists abundantly combined with lime. The process which he found to answer best is the following: To the juice of the house-leek add acetate of lime as long as any precipitate takes place. Wash the precipitate, and decompose it by means of diluted sulphuric acid in the manner directed by Scheele †.

Malic acid may be formed also by the action of nitric acid on sugar. If nitric acid be distilled with an equal quantity of sugar, till the mixture assumes a brown colour (which is a sign that all the nitric acid has been extracted from it), this substance will be found of an acid taste; and after all the oxalic acid which may have been formed is separated by lime-water, there remains another acid, which may be obtained by the following process: Saturate it with lime, and filter the solution; then pour upon it a quantity of alcohol, and a coagulation takes place. This coagulum is the acid combined with lime. Separate it by filtration, and edul-

Chap. II.

[#] Swedish Trans. and Crell's A nals for 1785.

[†] Ann. de Com. IZZiv. 127.

corate it with fresh alcohol; then dissolve it in dispilled water, and pour in acetate of lead till no more precipitation ensues. The precipitate is the acid combined with lead, from which it may be separated by diluted sulphuric acid.

Bouillon La Grange has larely endeavoured to prove, that this acid is nothing else than the acetic combined with a peculiar vegetable matter. He has shown that acetic acid exists in the juice of apples, and thinks he has proved that nitric acid forms with sugar a peculiar extractive stuff which exists also in apples †. But his observations do not appear to me to prove the coincidence between acetic and malic acids, as they possess properties so very different from each other.

Properties.

2. Malic acid, thus obtained, is a liquid of a reddish brown colour and a very acid taste. When evaporated it becomes thick and viscid like a mucilage or syrup, but it does not crystallize. When exposed to a dry atmosphere in thin layers, it dries altogether, and assumes the appearance of varnish.

When heated in the open fire it becomes black, swells up, exhales an acrid fume, and leaves behind it a very voluminous coal. When distilled, the products are an acid water, a little carbureted hydrogen gas, and a large proportion of carbonic acid ‡.

- 3. It is very soluble in water. It gradually decomposes spontaneously, by undergoing a kind of fermentation in the vessels in which it is kept.
 - 4. Sulphuric acid chars it, and nitric acid converts it

^{*} Savelish Truns. and Crel 's Annals for 1785.

[†] Gehien's Jour. 2d Seres. n. 173.

‡ Fourcroy, vii. 199.

into oxalic acid *. Hence it is evident that it is com- Chap. II. posed of oxygen, hydrogen, and carbon, though the proportions of these substances have not been ascertained.

Malic acid combines with alkalies, earths, and metallic oxides, and forms salts known by the name of malates.

- T is acid bears a strong resemblance to the citric, but differs from it in the tollowing particulars:
- 1. The citric acid shoots into fine crystals, but this acid does not crystallize.
- 2. The salt formed from the c tric acid with lime is almost insoluble in boiling water; whereas the salt made with malic acid and the same basis is readily soluble by boiling water.
- 3. Malic acid precipitates mercury, lead, and silver, from the nitrous acid, and also the solution of gold when diluted with water; whereas citric acid does not alter any of these solutions.
- 4. Malic acid seems to have a less affinity than citric acid for lime; for when a solution of lime in the former acid is boiled one minute with a salt formed from volatile alkali and citric acid, a decomposition takes place, and the latter acid combines with the lime and is precipitated.

^{*} Schoole and Hermhstadt.

SECT. XVI.

OF SUBERIC ACID.

Flistory.

Cork, a substance too well known to require any description, is the bark of a tree which bears the same name. By means of nitric acid, Brugnatelli converted it into an acid *, which has been called the suberic acid, from suber, the Latin name of the cork tree. Several chemists affirmed that this acid was the oxalic, because it possesses several properties in common with it. These assertions induced Bouillon La Grange to undertake a set of experiments on suberic acid. These experiments, which have been published in the 23d volume of the Annales de Chimie, completely establish the peculiar nature of suberic cid, by showing that it possesses properties different from those of any other acid.

Preparation. 1. Substic acid may be formed by pouring six parts of nitric acid of the specific gravity 1.261 on one part cork grated down, or simply broken down into small pieces, and distilling the mixture with a gentle heat as long as red vapours continue to escape. As the distillation advances, a yellow matter like wax makes its appearance on the surface of the liquid. While the matter contained in the retort is hot, it is to be poured into a glass vessel, placed upon a sand-bath over a gentle fire, and constantly stirred with a glass rod. By this

^{*} Creli's Annais, 1, 37.

lans it becomes gradually thick. As soon as white pours, exciting a tickling in the throat, begin to engage themselves, the vessel is removed from the th, and the mass continually stirred till it is almost dd.

By this means an orange-coloured mass is obtained the consistence of honey, of a strong and sharp odour He hot, but having a peculiar aromatic smell when

On this mass twice its weight of boiling water is to poured, and heat applied till it becomes liquid; and that part of it which is insoluble in water is to be parated by filtration. The filtered liquor becomes addy; on cooling it deposits a powdery sediment, a thin pellicle forms on its surface. The sediment be separated by filtration, and the liquor reduced a dry mass by evaporating in a gentle heat. This 18 18 suberic acid. It is still a little coloured, owing some accidental mixture, from which it may be pued either by saturating it with potash and precipitay it by means of an acid, or by boiling it along with rcoal powder.

. Suberic acid thus obtained is not crystallizable, Properties. when precipitated from potash by an acid it asmes the form of a powder; when obtained by evapotion it forms thin irregular pellicles.

3. Its taste is acid and slightly bitter; and when diswed in a small quantity of boiling water it acts upon throat, and excites coughing.

It reddens vegetable blues; and when dropt into solution of indigo in sulphuric acid (liquid blue, as it called in this country), it changes the colour of the Mution, and renders it green.

Chap. H.

4. Water at the temperature of 60° or even 70° dissolves only 17.8 part of its weight of suberic acid; and if the acid be very pure, only 11.2 th part: boiling water, on the contrary, dissolves half its weight of it.

When exposed to the air, it attracts moisture, especially it it be impure.

5. When exposed to the light of day, it becomes at last brown; and this effect is produced much somer by the direct rays of the sun.

When heated in a matrass, the acid sublimes, and the inside of the glass is surrounded with zones of different colours. If the sublimation be stopt at the proper time, the acid is obtained on the sides of the vessel in small points formed of concentric circles. When exposed to the heat of the blow-pipe on a spoon of platinum, it first melts, then becomes pulverulent, and at last sublimes entirely with a smell resembling that of distilled oil.

It is not altered by oxygen gas:—the other acids do not dissolve it completely. Alcohol developes an aromatic odour, and an ether may be obtained by means of this acid.

It converts the blue colour of nitrate of copper to a green; the sulphate of copper also to a green; green sulphate of iron to a deep yellow; and sulphate of zinc to a golden yellow.

It has no action either on platinum, gold, or nickel; but it oxidizes silver, mercury, copper, lead, tin, iron, bismuth, arsenic, cobalt, zinc, antimony, manganese, and molybdenum.

6. With alkalies, earths, and metallic oxides, it forms compounds known by the name of suberates.

SECT. XVII.

OF FORMIC ACID.

s acid is first mentioned in the Philosophical History. sactions for 1671, in a paper by Mr Ray, giving count of the observations of Mr Halse, and the exneuts of Mr Fisher, on the acid juice which is sponusly given out by ants, and which they yield when Mr Fisher compares this liquor with vinebut points out some differences between them. ely any addition was made to these facts till Marpublished a dissertation on the subject in the Ber-Iemoirs for 1749, in which he describes the meof obtaining the formic acid from the formica rur red ant, and points out its properties with his precision and method †. A new dissertation was ished on the same subject by Messrs Arvidson and n in 1782, in which the discoveries of Margraff confirmed, and many new particulars added. abstadt's paper on the same subject appeared in i's Annals for 1784. His researches were directed ly to the purification of the formic acid. strated that the juice of ants contained several fobodies, and among others, that a portion of malic might be detected in it. Richter published expents on formic acid about the year 1793, pointing

out a method of procuring it in a very concentrated **y** state *. Deveux soon after examined it, and found it 1: analogous to the acetic acid +. This common of the (.) French chemist was confirmed in 1802 by Fourcroy and 1783 Vauquelin who published a dissertation on ants, and r.... corcluded from their experiments, that the formical K:3. is nothing else than a mixture of the acetic and malic と予 The opinion maintain a by these celebraed * : philosophers induced Suersen to a samine the subjectly experiment. This chemist, in an elaborate dissertation on turnic acid, published in 180: \$, shows that most of the facts pointed out by Fourcroy and Vauquelin had been already ascertained by preceding chemists; that the experiments which they detail were not sufficiented warrant their conclusions; that formic acid when properly prepared contains no malic acid; and that it possesses properties different from the acetic. These conclusions have been farther confirmed by the experiments of Gehieu ||.

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Prepara-

The simplest method of procuring formic acid in a state of purity is that of Margraff as corrected by Richter. Suersen accordingly had recourse to it. This method is as follows: Infuse any quantity of ants in about thrice their weight of water, put the mixture into a silver or tin coppered still, and draw off the water by distillation as long as it continues to come over without any burnt smell: for the distillation must be stop-

[#] Gehlen, iv. 7.

[†] Fourctoy, x. 491. Eng. trans. I have not seen either the disentation of Richter of Deyeux.

¹ P. . M., v r 8.

[§] Gehlen's Johr. iv. 1.

¹ Ain. de Chim. lx. 78.

is soon as that smell begins to be perceived. Sate- Chap. II. the water in the receiver with carbonate of potash, and torate to dryness. Mix the white mass thus obwith as much sulphuric acid, previously diluted its weight of water, as is sufficient to saturate the b. Introduce the mixture into a retort, and distil ly to dryness. The liquid which comes over into receiver is to be again rectified by a very moderate ato get rid of any portion of sulphuric acid that be present. It is now pure formic acid.

formic acid thus procured is colourless like water. Proportion anot contain make acid, because that acid cannot be Med over along with water.

smell is peculiar, and very different from that of ahacid. Its taste i acid. I reide is vegetable blues. is specific gravity varies from 1.100 to 1.113; mean the most concentrated acetic acid is only 1.080. does not appear susceptible of being brought to the of crystals; at least Lowitz attempted in vain to bit it in that state, though he succeeded readily with ie acid.

Notwithstanding its greater specific gravity, it is cae of neutralizing much less of alkaline bodies than ic acid, as appears from the following trials of men. He reduced formic and acetic acids, each to apecific gravity 1.0525, and ascertained how much bonate of potash, carbonate of lime, and carbonate of enesia, the same weight of each of these liquids was able of neutralizing. The result may be seen in the lowing little Table.

> Fermic Acid. L'me 166 231 Magnesia 150 213

Analogy with acetic acid.

Such are the differences between the formic and acetic acids pointed out by Suerson. In other respects there appears to be a striking analogy between the two acids,

Both of them form soluble compounds with all the alkalies and alkaline earths. They precipitate no metal from its solution, and of course form likewise soluble compounds with all the metallic oxides; but there is one striking difference pointed out long ago by Mr Fisher, which has not been noticed by any modern chemist. It is this: Acetate of lead when distilled yields no acid liquor whatever; but formic acid may be obtained from formate of lead precisely as acetic acid is procured from acetate of copper.

Upon the whole, though the experiments of the German chemists are sufficient to separate the formic and acetic acids, their analogy in other respects is so great, that farther researches are still requisite to elucidate the subject: Above all, an examination of the salts which the formic acid forms with the different bases ought to be undertaken, in order to ascertain whether formic acid may not be a combination of acetic acid with some unknown body. Gehlen indeed affirms that the formates differ from the acetates, and instances the formate of copper, which he says is a blue salt crystallizing in cubes.

SECT. XVIII.

GENERAL REMARKS.

1. The preceding Sections of this Chapter contain an

secount of all the acids at present known, except five; which for obvious reasons have been referred to other Acids emis parts of this Work. The first three of these will be described in the following Chapter. The other two, namely, the rosacic and amniotic, are never employed as instruments of analysis. Their acid characters are ambiguous, and their properties have been but imperfectly investigated. It appeared better, therefore, to reserve them till we came to treat of those animal substances in which they are found.

2. Several acids usually described by chemists will Acids denot be found in the preceding Sections, because their peculiar nature has been destroyed by the more precise investigation of modern analysis. The most remark. able of these acids are the five following.

I. LACTIC ACID.

When milk is kept for some time it turns sour. Scheele examined the acid thus evolved, and gave it the name of lactic acid. He directs the following method of procuring it.

Evaporate a quantity of sour whey to an eighth Preparapart, and then filtrate it: this separates the cheesy part. Saturate the liquid with lime-water, and the phosphate of lime precipitates. Filtrate again, and dilute the liquid with three times its own bulk of water; then let fall into it oxalic acid, drop by drop, to precipitate the lime which it has dissolved from the lime-water; then add a very small quantity of limewater, to see whether too much oxalic acid has been added. If there has, oxalate of lime immediately pre-Evaporate the solution to the consistence of cipitates.

honey, pour in a sufficient quantity of alcohol, and his trate again; the acid passes through dissolved in the alcohol, but the sugar of milk and every other substance remains behind. Add to the solution a small quantity of water, and distil with a small heat; the alcohol passes over and leaves behind the lactic acid dissolved in water *.

From the properties of the acid procured by this process, Scheele concluded that it was very analogous to the acetic, but that it wanted something to bring it that state. He even pointed out a method of procuring vinegar from milk in considerable quantity, and ascertained that it always yielded a little when distilled †.

From the recent experiments of Bouillon La Grange, we learn that the lactic acid is not a peculiar acid, but a mixture of the following substances: 1. Acetic acid; 2. Muriate of potash; 3. A little iron; and 4. An animal matter ‡.

II. ZOONIC ACID.

WHEN animal substances and those vegetable principles which possess similar properties are distilled, the product contains an acid liquid, which Berthollet considered as peculiar, and called zoonic acid.

To obtain it, he mixed lime with the distilled liquid, evaporated nearly to dryness, then diluted and filtered.

[•] Scheele, Stockbolm Trans. 1780.

[†] Scheele, ii. 66.

¹ Ann. de Chim. L 288.

horic acid was then added, and the mixture dis-The receiver contained the zoonic acid *. Chap. II.

m the more recent experiments of Thenard, we that this supposed acid is no other than the acelding in solution a peculiar animal matter resemoil †.

III. PYROMUCOUS ACID.

ed, among other products there is always a notacantity of an acid liquid. This acid, when rectibtained the names of syrupous acid, and afterwards acous acid. It is now known from the recent exents of Fourcroy and Vauquelin, that this acid is ag else than the acetic, holding in solution a porf empyreumatic oil ‡.

L V. Pyrolignous and Pyrotartarous Acids.

more or less of an acid juice: the same remark is to the salt called tartar. These liquids were guished by the name of pyrolignous and pyrotartacids: but they are now known to be only the aces guised by the presence of a peculiar oil §. It is r to observe, however, that Gehlen has lately sed that the last of these acids is quite different

[#] Ann. de Chim. xxvi. 86.

^{† 1}bid. zliii 276.

[‡] Ibid. xxxv. 131.

Fourcroy and Vauquelin, Ann. de. Chim. xxxv. 131.

from the acetic. When cream of tartar is distilled, if yields an acid liquor, which, by proper concentration deposites brownish crystals constituting an acid which cannot be acetic, and which differs equally from the last taric.

- which remain to be described, amount to 34. Of there one, the muriatic, possesses the characters of a simple substance; seven are products of combustion, in nine supporters of combustion. The remaining reare combustible. Oxygen is essential to the acidity of the products and supporters. The combustible and almost all contain it; but it cannot be shown to be a sential to their acidity. It has been usual with the mists to divide the combustible acids into vegetable and animal, because most of them are obtained from the vegetable and animal kingdoms. Several of the other acids have been denominated mineral for a similar resort.
- 4. Let us take a general view of all the acids, in of der to ascertain in what degree each of them possess those properties which are usually considered as characterising acid bodies. These properties are, 1. As acid taste; 2. The power of convening vegetable blue into red; 3. Solubility in water; 4. The property of forming salts.

Tauce of the acids.

All the acids have a sour taste, except the exymumtic, the tungstic, the columbic, and the uric. In seven the acidity is mixed, and even almost concealed, by some other more powerful taste. But this will be best understood by inspecting the following TABLE.

[·] Ann de Chim. la 79.

de.	Taste.	Acida.	Taste.	Chap. II.
ic abic	None	Sulphuric Phosphoric Phosphorous Nitric		
ic	Hot, acrid	Muriatic Fluoric Acetic Oxalic	Acid, strong	
uriatic	Astringent			
ic tylic	Acid, as- tringent	Tartaric Citric Kinic		
ic pdic pic	Acid, me- tallic	Malic		
ic horic ic	Acid, bitter	Mellitic	Acid, weak	,
nrous	Acid, sulph.			<u> </u>

All the acids are destitute of smell excepting the ing:

Acide.	Smell.	
ulphurous	Sulphureous	
'hosphorous	Phosphureted hydrogen	
Auriatic.		
luoric	Muriatic acid	
)xymuriatic	Aqua regia	
Iyperoxymuriatic	Nitrous gas	

Odour.

Acids.	Smell.	
Nitric		
Acetic	Vinegar	
Succinic *	Distilled nutgalls	
Camphoric	Saffron	
Benzoic* Moroxylic	Benzoin	

Action on vegetable blues.

Carbonic acid, though a gas, has no smell whatever.

6. All the acids convert vegetable blues to red; but they differ exceedingly in the extent to which they possess this property. The greater number convert & most all vegetable blues (except indigo) to red; while some, the carbonic, for instance, only act upon the most delicate, as the tineture of turnsol. The cause of this change of colour has not been ascertained. cases it seems to be in consequence of the saturation of an alkaline or earthy substance which was previously combined with the colouring matter. This is the case If this held always, evidently with respect to litmus. the conversion of vegetable blues into red would be merely in consequence of the affinities between acids and alkaline bodies, and of the superior affinity which they have for them.

Solubility in water.

7. All the acids are soluble in water; but in this respect also they differ exceedingly from each other. Two of them are so soluble in water, that they have never

^{*} In these two the odour is accidental, and owing to an oil.

n obtained except in combination with that liquid. Chap. II. see are,

- 1. Sulphuric,
- 2. Nitric.

a solid or crystallized form. Six of them are es, which are readily absorbed by water to a certain ount; after which the water being saturated, refuses absorb any more. The weight of each absorbed by parts of water may be seen in the following TABLE.

- 1. Muriatic.....54.8
- 2. Sulphurous..... 9.15
- 3. Carbonic..... 0.18
- 4. Oxymuriatic.....unknown
- 5. Hyperoxymuriatic.....unknown
- 6. Fluoric.....unknown

The weight of the remaining acids, when solid or stallized, soluble in 100 parts of water, is exhibited ac following TABLE.

Phosphoric Phosphorous Very soluble, proportion not determined. Acetic Tartaric Malic Oxalic..... 50 Boracic..... Saclactic..... Succinic..... 1.0 Suberic..... 0.7 Camphoric..... 0.5

Book IL Division 11. Benzoic..... 0'2
Molybdic 0'1

All the acids are more or less soluble in alcohol, as cept phosphoric acid and the metallic acids. The phuric, natric, and oxymuriatic, as we shall see the wards, have the property of decomposing alcohol.

8. All the acids are capable of combining with the lies, earths, and metallic oxides, and of forming the them compounds known by the name of salts. It substance which wants this property ought to be made as an acid.

Combination with alkalies and cartha.

Composi-

9. The two first classes of acids, as far as is know at present, are more simple than the third; as the sol belonging to them contain only two ingredients ad whereas the combustible acids contain three too four constituent parts. These last are or imposed under of oxygen, hydrogen, and carbon. Hence the inson that several of them may be converted into other by the action of nitric acid. This acid alters the proportion of their ingredients, either by abstracting put of their carbon and hydrogen, or by communicator oxygen, or by both together. But it is impossible to explain precisely what these changes actually amount to, without being acquainted with the component pans of every combustible acid, the manner in which these component parts are combined, and the affinities which exist between each of them. This, however, is very far from being the case at present. Though a wast number of experiments have been made on purpose to throw light on this very point, the difficulties which were to he encountered have been so great, that no accurate results have yet been obtained. All that is known at present is an approximation towards the component ree acids. This may be seen in the follow- Chap. II4

Acids.	Ozygen.	Carbon.	Hydrogen.
cetic	50-19	35.87	11.94
xalic	64	32	4
artaric	70.5	18	10.2

cids was composed by chemists, tha each cids was composed of a peculiar compound dical, combined with oxygen. The radical sed to be a combination of carbon and hydrono proof whatever has been given of the is opinion, nor has the smallest evidence been that any such radicals exist. It is much more hat all the ingredients of the acids are combined ther, and constitute a triple combination.

CHAP. III.

OF COLORIFIC ACIDS.

UNDER the name of colorific acids (till some more appropriate appellation occur), I include three substances, which possess such analogous properties that they ought to be classed together. As chemical bodies, they are all applied to nearly the same purposes; namely, to detect the presence of metallic bodies, and to separate them from other substances. They act with great energy upon the metallic solutions, and at the same time precipitate them in powders remarkable for the intensity and variety of their colours. It was this circumstance that induced me to apply to them the epithet colorific.

Two of these bodies, namely, prussic and gallic acids, have been long considered as acids by chemists; though the second does not seem capable of neutralizing alkalies, and therefore is scarcely entitled to the name. A third of them, sulphureted bydrogen, has been shown by Berthollet to act as an acid; but the compounds which it forms with alkalies are only transient. Its importance depends upon the energy with which it acts upon metallic bodies. Along with these three acids I class a fourth substance, tannin, which, though not an acid, is applied to the same purposes as the other

three. Indeed it would be difficult to assign any reason for giving the title of acid to gallic acid, which would exclude tannin. Upon the whole, perhaps, it would be better to distinguish these bodies altogether from the acids, and to assign them a peculiar name of their own. In describing them, it may be proper to begin with the body in which the acid characters are best marked, and to terminate with that in which they can no longer be perceived. With this view they may be arranged in the following order:

Chap. III.

- 1. Sulphureted hydrogen
- 2. Prussic acid
- 3. Gallic acid
- 4. Tannin.

These bodies may be distinguished by the following characters:

- 1. They unite with alkaline bodies, but are all incapable of neutralizing them except the first; and the compound which it forms cannot be exposed to the air without decomposition.
- 2. They act with great energy upon metallic solutions, usually entering into combination with the oxide, and precipitating it in the state of an insoluble powder.
- 3. They have a tendency to enter into triple compounds with a variety of bodies, especially metallic oxides and alkalies; except the sulphureted hydrogen, which is commonly decomposed by metallic solutions.

Buck fi. 1) vision II.

SECT. I.

OF SULPHURETED HYDROGEN.

This substance forms the link by which the colorife acids are joined to the acids strictly so called. The Germans have given it the name of bydrothionic acid. The method of preparing this substance, and the most remarkable of its properties, have been detailed in a preceding part of this Work. As an instrument of chemical analysis it is usually employed in two states.

1. Dissolved in water; in which state it is called liquid sulphureted bydrogen.

2. Combined with alkalies, by causing a current of sulphureted hydrogen gas to pass through an alkaline solution till the liquid refuses to absorb any more. The liquid is then heated, to expel the excess of gas. In that state the compound is called an alkaline bydrosulphuret.

SECT. II.

OF PRUSSIC ACID.

We are indebted to an accident for our knowledge of this important substance. About the year 1710 Dies-

[•] See Vol. L. p. 89.

bach, a preparer of colours in Berlin, wishing to prepare some lake by precipitating a decoction of cochineal, alum, and green vitriol, with potash, borrowed some alkali for that purpose from Dippel. This chemist was the discoverer of a peculiar animal oil which goes by his name. He prepared it from blood; and the ikali with which he furnished Diesbach had been employed in the process. Instead of the red precipitate which he expected, a beautiful blue powder fell to the bottom. On mentioning the circumstance to Dippel, that chemist ascribed the formation of the powder to the action of his alkali on the alum and vitriol. It is not unlikely that he had calcined the potash together with a portion of blood. Be that as it may, he succeeded in discovering a method of procuring the blue powder at pleasure, and it was announced as a pigment in the Berlin Miscellanies for 1710. The preceding history, however, was only communicated to the public by Stahl 20 years after ".

This powder was called Prussian blue; and the method of procuring it remained concealed, because it had become a lucrative article of commerce, till Dr Woodward published a process in the Philosophical Transactions for 1724, which he had procured, as he informs us, from one of his friends in Germany. This method was as follows: Detonate together four ounces Preparaof pitre and as much tarter, in order to procure an extemporaneous alkali; then add four ounces of dried bullock's blood; mix the ingredients well together, and put them into a crucible covered with a lid, in which

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Stahl's Experimente, Observat. Animadvers. ecc. numero. Chim. et Phys. p. 281.

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there is a small hole; calcine with a moderate fire till the blood emits no more smoke or flame capable of blackening any white body exposed to it; increase the fire towards the end, so that the whole matter contained in the crucible shall be moderately but sensibly red. In this state throw it into four pounds of water, and boil it for half an hour. Decant off this water, and continue to pour on more till it come off insipid. Add all these liquids together, and boil them down to four pounds. Dissolve an ounce of sulphate of iron in half a pound of water, and eight ounces of alum in four pounds of boiling water: mix all the three solutions together while boiling hot. An effervescence takes place, and a powder is precipitated of a green colour. Separate this precipitate by filtration, and pour muriatic acid upon it till it becomes of a beautiful blue; then wash it with water and dry it *.

Different explanations were given of the nature of this precipitate by different chemists. Mr Brown immediately repeated the process of Woodward, ascertained that other animal substances, as beef, may be substituted for blood; that the alum is useful only to dilute the colour; and that the blue pigment is produced by the action of the alkali (altered by blood) on the iron of the vitriol. He ascertained, too, that prussian blue is insoluble in muriatic acid, and that the green colour is owing to a mixture of prussian blue and oxide of iron, and that the muriatic acid developes the blue colour by dissolving the oxide of iron +.

These facts were of considerable importance; but they

[#] Phil Trans. xxxiii. 15. † Ibid. 1724. xxxiii. 17.

threw no light upon the theory of the process. An ex- Chap. Hi. planation of this was first attempted by Geoffroy, who bad ascertained that any animal body whatever might be substituted for blood. According to him, the blood communicates a portion of inflammable matter, or phlogiston, to the alkali, and this inflammable matter revives the iron of the vitriol and brings it to the metallic state. A greater quantity of blood, by increasing the inflammable matter, will enable the alkali to revive still more of the iron, and thus to strike a blue at once, instead of a green . Though this explanation was approved of at the time by the best chemists, it was far from satisfactory t. Macquer soon after proceeded, by way of experiment, and added a new step to the facts ascertainad by Brown.

That celebrated chemist ascertained the following Experi-facts: 1. When an alkali is added to a solution of iron Macquein any zeid, the iron is precipitated of a yellow colour, and soluble in acids; but if iron be precipitated from an acid by an alkali prepared by calcination with blood (which has been called a prussian alkali), it is of a green colour. 2. Acids dissolve only a part of this precipitate, and leave behind an insoluble powder which is of an intense blue colour. The green precipitate therefore is composed of two different substances, one of which is prussian blue. 3. The other is the brown or yellow oxide of iron; and the green colour is owing to the mixture of the blue and yellow substances. 4. When heat is applied to this prussian blue, its blue co-Jour is destroyed, and it becomes exactly similar to com-

Brok II, Division II. mon oxide of iron. It is composed therefore of iron and some other substance, which heat has the property of driving off. 5. If it be boiled with a pure alkal. it loses its blue colour also, and at the same time the al. kali acquires the property of precipitating of a blue colour solutions of iron in acids, or it has become precisely the same with the prussian alkali. 6. Prussian blue, therefore, is composed of iron and something which a pure alkali can separate from it, something which has a greater affinity for alkali than for iron, 7. By boiling a quantity of alkali with prussian blue, it may be completely saturated with this something, which may be called colouring matter, and then possesses the properties of a neutral salt. 8. No acid can separate this colouring matter from iron after it is once united with it. 9. When from dissolved in an acid is mixed with an alkali saturated with the colouring matter, a double d composition takes place; the acid unites with the alkali, and the colouring matter with the iron, and forms prussian blue. 10. The reason that, in the common method of preparing prussian blue, a quantity of yellow oxide is precipitated, is, that there is not a sufficient quantity of colouring matter (for the alkali is never saturated with it) to saturate all the iron displaced by the alkali; a part of it therefore is mixed with prussian blue. Muriatic acid dissolves this oxide, carries it off, and leaves the blue in a state of purity. - Such were the conclusions which Macquer drew from his experiments; experiments which not only discovered the composition of prussian blue, but threw a ray of light on the nature of affinities, which has contributed much towards the advancement of that important branch of chemistry.

The nature of the colouring matter, however, was still unknown. Macquer supposed it to be phlogiston. According to him, prussian blue is nothing else than iron supersaturated with phlogiston. This overdose protects the iron from acids, and prevents the magnet from acting on it. Heat drives off this dose, and leaves the prussian blue in the state of common iron. From this theory, which differed but little from that of Geoffroy, the alkali saturated with the colouring matter of prussian blue received the name of phlogisticated alkali. Macquer having observed that it did not act on alkaline and earthy solutions, while it precipitated all the metals,

proposed it as an excellent test for detecting the pre-

sence of these last bodies.

The subsequent experiments of chemists threw an air of suspicion on Macquer's theory. Baumé ascertained, that when prussian blue is distilled, it always vields a portion of animal oil +; a product not very likely to appear if the powder contained nothing but philogiston and iron. Deyeux and Parmentier, Bergman, Erxleben, Delius, and Scopoli, submitted p. ussian blue to distillation, and obtained a quantity of ammonia. Fontana ascertained that prussian blue detonated with nitre. Landrians obtained, by distillation, a little acid liquid and oil, and a great quantity of azotic gas and carburcted hydrogen gas. These facts were still more inconsistent, if possible, with Macquer's theory. Morveau advanced another in 1772; namely, that the phlogisticated alkali, besides phlogiston, contained also an acid which acted the principal part in the phe-

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See Macquer's Dictionary, i 177. | † Baume's Chemitty, ii box.

nomena produced*. Sage affirmed that the colouring matter in phlogisticated alkali was phosphoric acid; but this opinion was refuted by Lavoisier †. Bergman also announced his suspicions that it was an acid, but an unknown one 1.

Such was the knowledge of chemists respecting the nature of this colouring matter, when Scheele all at once removed the veil, and explained its properties and composition. This he performed in two dissertations on prussian blue, published in the Stockholm Transactions for 1782 and 1783 ||.

He observed that the prussian alkali, after being exposed for some time to the air, lost the property of forming prussian blue; the colouring matter must therefore have left it.

Discovery of prussic acid.

He put a small quantity of it into a large glass globe, corked it up, and kept it some time; but no change was produced either in the air or the prussian alkali. Something must therefore displace the colouring matter when the alkali is exposed to the open air, which is not present in a glass vessel. Was it carbonic acid gas? To ascertain this, he put a quantity of prussian alkali into a glass globe filled with that gas, and in 24 hours the alkali was incapable of producing prussian blue. It is therefore carbonic acid gas which displaces the colouring matter. He repeated this experiment with this difference, that he hung in the globe a bit of paper which had been previously dipped into a solution of sulphate of iron, and on which he had let fall two drops

^{*} Digressions Academiques, p. 24%.

⁺ Mem. Par. 1777, p. 77.

Notes on Scheffer, ; 165.

[&]quot;Scheele, ii. 141.

of an alkaline lixivium in order to precipitate the iron. This paper was taken out in two hours, and became cowered with a fine blue on adding a little muriatic acid. Carbonic acid, then, has the property of separating the colouring matter from alkali without decomposing it.

He found also that other acids produce the same efbect. Hence he concluded that the colouring matter might be obtained in a separate state. Accordingly he made a great many attempts to procure it in that state, and at last hit upon the following method, which sucteeds perfectly.

Mix together ten parts of prussian blue in powder, Preparafive parts of the red oxide of mercury, and thirty parts of water, and boil the mixture for some minutes in a plass vessel. The blue colour disappears, and the mixture becomes yellowish green. Pour it upon a filter; and after all the liquid part has passed, pour ten parts of hot water through the filter to wash the residuum completely. The oxide of mercury decomposes prussian blue, separates its colouring matter, and forms with it a salt soluble in water. The liquid therefore which has passed through the filter contains the colouring matter combined with mercury. The other component parts of the prussian blue being insoluble, do not pass through the filter. Pour this mercurial liquid upon 21 parts of clean iron filings, quite free from rust. Add at the same time one part of concentrated sulphyric seid, and shake the mixture. The iron filings are dissolved, and the mercury formerly held in solution is precipitated in the metallic state. The cause of this sudden change is obvious: The iron deoxidizes the mercury, and is at the same instant dissolved by the miphuric acid, which has a stronger affinity for it than Vol. II.

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bided a quantity of sal ammoniae in small pieces, which he pushed to the bottom of the melted mixture, kept it in the fire for two minutes till it had ceased to give out vapours of ammonia, and then threw it into a quantity of water. The solution possessed all the properties of the prussian alkali. Thus Mr Scheele succeeded in forming the colouring matter.

This colouring matter was called prussic acid by Morveau in the first volume of the chemical part of the Encyclopedie Methodique; an appellation which is now generally received, and which therefore it will be proper to employ in the remaining part of this Section.

These admirable experiments of Scheele were repeatand carried still farther by Berthollet in 1787; who applied to the explanation of the composition of the colouring matter the light which had resulted from his previous experiments on the component parts of ammonia. This illustrious chemist, not inferior to Scheele in ingenuity and address, ascertained, in the first place, that the phlogisticated alkali is a triple salt, composed of prussic acid, the alkali, and oxide of iron; that it may be obtained in octahedral crystals; and that when mixed with sulphuric acid, and exposed to the light, it lets fall a precipitate of prussian blue. His next object was to ascertain the component parts of prossic acid. When oxymuriatic acid is poured into prussic acid, obtained by Scheele's process, it loses its oxygen, and is converted into common muriatic acid. At the same time the prussic acid becomes more odorous and more volatile, less ca, able of combining with alkalies, and precipitates it " from its solutions, not blue, but green. Thus prussic acid, by combining with exygen, acquires new properties, and is converted into a new substances

which may be called oxy-prussic acid. If more oxymuriatic acid gas be made to pass into prussic acid, and
it be exposed to the light, the prussic acid separates
from the water with which it was combined, and precipitates to the bottom in the form of an aromatic oil;
which heat converts into a vapour insoluble in water,
and incapable of combining with iron. When the
green precipitate, composed of oxy-prussic acid and
iron, is mixed with a pure fixed alkali, the oxy-prussic
acid is decomposed, and converted into carbonate of
ammonia.

From these experiments, Berthollet concluded, that prussic acid does not contain ammonia ready formed; but that it is a triple compound of carbon, hydrogen, and azote, in proportions which he was not able to accertain. This conclusion has been still farther verified by Mr Clouet, who found, that when ammoniacal gas is made to pass through a red hot porcelain tube containing charcoal, a quantity of prussic acid is formed. This experiment does not succeed unless a pretty strong heat be applied to the tube †.

Fourcroy and several other chemists believe, that the prussic acid contains also a portion of oxygen in its composition, testing chiefly upon an experiment of Vauquelin ‡. This is certainly possible, though it has not

^{*} Ann. de Chim. zi. 30. † Jour. de l' Ecole Polytechn. 1. iii. 436.

¹ Vauquelin's experiments were as follows:

EXPLR. I. Put into a retort 100 parts of the muriate of ammonia, 50 parts of time, and 24 parts of charcoal in fine powder; adapt to the retort a receiver containing a slight solution of the sulphate of iron, and immerse into it the beak of the retort; then apply a brisk heat, and continue the action of the fire until nothing more is disengaged.

been proved; and the experiments of Berthollet render Chap. II. at somewhat unlikely.

Having thus traced the gradual progress of philosophers, in ascertaining the nature of the prussic acid, it only remains to give an account of its properties, which were first examined by the indefatigable Scheele.

Prussic acid obtained by Scheele's process is a co. Properties. lourless liquid like water. It has a strong odour, resembling that of the flowers of the peach, or of bitter almonds. Its taste is sweetish, acrid, and hot, and apt to excite cough. It does not alter the colour of vegetable blues. It is exceedingly noxious when taken in- Poisonous. ternally, even exposure to the furnes of it proves fatal to small animals. The poisonous qualities of the distilled water of bitter almonds and leaves of laurel are ascribed at present to this acid, which is known to exist in these waters in considerable quantity.

It is very volatile, and evidently capable of assuming Gaseous. the gaseous form; though hitherto it has scarcely been examined in that state. The following experiment of

Expen. II. Put into a retort 100 parts of the muriate of animonia, 50 parts of semi-virreous oxide of lead, and 25 parts of charcoal; adapt a receiver containing a solution of sulphate of iron, and proceed as before. Sur well the liquors contained in the receivers, and expise them to the sar t r several days, in order that the combination between the oxide of from and the prusse acid may be perfect, and that the prussate of iron may absorb as much oxygen as is necessary for its passing to the state of blue pruss are, and for its being proof against acids: then pour into these liquors equal quantities of sulphuric acid well diluted with water, and you will have pressian blue, the quantities of which will be as one to are; that is to say, the prussian bise of the experiment in which Vauqueld emp eyed oxide of lead, was see three more abundant than that of the experiment in which he improved only his cito disengage the animonis,

Book II. Diva on II. Grindel, if accurate, shows that it may be procured a gas. He put a quantity of prussian blue into a glas flask, and fitting to it a bent tube, applied the heat of a lamp; as soon as the smell of bitter almonds became perceptible, the tube was plunged under mercury, and the gas evolved received in a jar containing liquid possess. The gas was absorbed by the potash, and the aquid, on being evaporated, deposited crystals, which possessed all the properties of prussiate of potash.

Not readily acted on by heat.

From the recent experiments of Richter and Bucholz, we learn that this acid is capable of resisting a much more violent heat without decomposition, than may of the vegetable acids. It is formed only when the potash as d blood are exposed to a red heat. When united to potash, a considerable heat may be applied to the combination without occasioning decomposition; but when the salt is dissolved in water prussic acid is immediately disengaged, as is evident by the odour of bitter almonds evolved; while at the same time a portion of ammonia and of carbonic acid is formed. Thus we see that water destroys the combination of this acid and alkalus. Buchoiz supposes that the effect is produced by the mutual action of the acid and water which decompose each other to These facts explain the reason of the facility with which the alkaline prussiales are decomposed by mere exposure to the air, and by all the acids, and show us that they can scarcely be applied to any useful purpose in chemistry, even if their difficult formation did not present an unsurmountable barto their introduction.

The presence of a metallic oxide serves to fix the con- Thap. Ht. stituents of prussic acid, and to prevent them from being sected upon by water or any other body. Hence these salts into which a metallic oxide enters are much more permanent in their nature, and of course may be used for chemical purposes.

Prussic acid then may be prepared as a reagent in four states: 1. In a state of purity, either gaseous or dissolved in water. 2. United to alkalies. 3. United so alkalies and metallic oxides at once in the state of a neutral salt. 4. United to metallic oxides alone.

15th, In the state of prussic acid its action on metallic Action co solutions is but feeble. Of all the metallic solutions tried by Scheele, pure prussic acid occasioned only a precipitate in three: namely,

1. Nitrate of silver precipitated white,

2. Nitrate of mercury..... black.

3. Carbonate of iron..... green becoming blue. It has no action on the oxides of

1. Platinum. 4. Lead.

7. Manganese,

2. Iron. 5. Bismuth, 8. Arsenic.

3. Tie, 6. Antimony, 9. Molybdenum.

- 10. Gold precipitated by the alkaline carbonates is rendered white by this acid.
- 11. It disengages carbonic acid from the oxide of ariver, precipitated by the same alkalies; but the oxide semains white.
- 12. It dissolves red oxide of mercury, and forms with it a salt which may be obtained in crystals.
- 13. Oxide of copper precipitated by carbonate of potash effervesces in it, and acquires a slight orange-yellow colour.
 - 14. Oxide of iron precipitated from the sulphate of

Book II. Davaion II. iron by carbonate of potash, effervesces in it, and becomes blue.

15. Oxide of cobalt precipitated by the same alkalı, gives in it some marks of effervescence, and becomes yellowish brown *.

2d, From the experiments of Scheele, Richter, and Bucholz, we learn, that the alkaline prussiates act with sufficient force upon metallic solutions, and produce more beautiful precipitates than the triple prussiates; but, for the reasons already assigned, they cannot be introduced into use with advantage.

digesting alkaline bodies on prussian blue, the oxide of iron is the metallic body which enters into combination with the acid and base, and constitutes the compound a triple sait. Potash has, with one consent, been adopted by chemists as most convenient; but other alkaline bodies would doubtless answer equally well. Ferruginous prussiate of potash, then, is the substance usually employed as a reagent. It is a yellow-coloured salt which crystallizes in flat cubes. It is used to detect the presence of metallic bodies by the colour of the precipitate formed, and in an especial manner to detect iron, which it does by the blue colour that the solution assumes; and to free solutions from iron, which it does by precipitating the iron in the form of prussian blue.

Ferraginous prusstate of potseh-

Prumiate of mercury.

4th, The only soluble combination of prussic acid and a metallic oxide is prussiate of mercury: the method of obtaining which is described above. It is a white salt of a disagreeable metallic taste, crystallizes in

^{*} Stheek, 11. 169.

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needles, and forms a colourless solution in water. This salt is employed with great advantage in several cases. By means of it, for example, Dr Wollaston has pointed out a method of separating palladium with facility from crude platina. Dissolve crude platina in nitro-muriatic acid; throw down the platinum with sal ammoniac, neutralize the remaining solution with an alkali, and then drop in prussiate of mercury; a yellow precipitate gradually forms, which, when heated to redness, leaves pure palladium.

The only earthy bodies precipitated by the prussiates are zirconia and yttria. This property distinguishes these two earths from all the rest, and points out an analogy between them and the metallic oxides.

SECT. III.

OF GALLIC ACID.

HERE is an excrescence, known by the name of nut- History gall, which grows on some species of oaks. This substance contains a peculiar acid, called from that circumstancogallic acid; the properties of which were first examined with attention by the commissioners of the Academy of Dijon, and the result of their experiments was published in 1777, in the third volume of their Elements of Chemistry *. In these experiments, however, they employed the infusion of galls, in which the

* Vol. III. p. 493.

Book II. Division II. Pr paraacid is combined with tannin. It was reserved in Scheele to obtain it nearly in a state of purity.

eold water, a sediment, which proved on examination to have a crystalline form and an acid raste. By letting an infusion of galls remain a long time exposed to the air, and removing now and then the mouldy skip which formed on its surface, a large quantity of the sediment was obtained; which being edulcorated with cold water, redissolved in hot water, filtrated and exporated very slowly, yielded an acid salt in crystals as fine as sand.

Deyeux has proposed a much speedier method of obtaining gallic acid †; but it does not succeed without a good deal of precaution. It consists in exposing pounded nut-galls in a large glass retort to a heat castiously and slowly raised. A number of brilliant white crystalline plates are amblimed, which possess all the properties of gallic acid. Care must be taken not to apply too great a heat, and to stop the process before any oil begins to come over, otherwise the crystals will be redissolved, and the whole labour lost.

Mr Davy has lately pointed out another method which yields gallic acid in a state of considerable parity. Boil for some time a mixture of carbonate of barytes and infusion of nut-galls. A bluish green liquid is obtained, which consists of a solution of gallic acid

Stubbelm Trans. 1786.—The crystals obtained by this method shows contain a portion of tannin, and are of a brown colour.

[†] This method was in fact discovered by Schoole; but Deyout repeated it, and pointed out the proper precautions.—See Crell's Anali; i 29. Eng. Transl.

and barytes. Filter and saturate with diluted sulphuric acid. Sulphate of barytes is deposited in the state of an insoluble powder, and a colourless solution of gal-Arc acid remains behind .

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Many other processes besides these have been proposed by different chemists, but as they are all liable to considerable objections, it is not necessary to enumerate them. The following method proposed by Richster, though expensive and tedious, is one of the best :

" Infuse in cold water one pound and a half of gall Method of nuts, previously reduced to fine powder, taking care frequently to agitate the mixture. Pass the liquid through a cloth; add water to the pulp which refuses to go birough, and again put it through the cloth, using a press to separate the water. Join the liquors, and with a gentle heat evaporate them, and a matter of a dark brown colour, and very brittle, will be obtained.

" Pure alcohol poured on this matter, reduced to a fine powder, acquires a pale straw colour. The deposite infused again in alcohol communicates but little colour to it. The brown residuum now left is composed almost wholly of pure tannin. Mix the two alcoholic extracts, which distil in a small retort to one eighth. What remains will be almost a solid mass. Pour water on it, and expose it to a gentle heat, and you will obtain a clear and almost colourless solution.

" Evaporate this solution, and you will obtain from it very small, white, prismatic crystals. The liquor furnishes more, but they are commonly a little coloured. It is sufficient to levigate them with water to ob

[·] Journal of the Royal Inchit. L. 274.

of crystals is procured from one pound of galls; there crystals are extremely light, and consequently occupy considerable space *."

Method of Proust.

The method of Scheele is by far the cheapest; but it never yields a pute acid. Mr Proust has proposed the following method of remedying this defect: Forma strong infusion of galls, set it aside till impure crystals of gallic acid are deposited. Dissolve these crystals in water, and drop muriate of tin cautiously into the solution. Flocculi are deposited, and the solution becomes clear. Filter and evaporate. Pure crystals of gallic acid are deposited, which require only to be dried upon blotting paper +. I have tried this method of Proust, but it did not succeed. Berthollet has proposed as a substitute to heat this solution of gallic acid with recently precipitated oxide of tin. But this method likewise failed in the hands of Bouillon La Grange, who has lately published a dissertation on gallic acid, and endeavoured to prove that it is merely acetic acid combined with tannin and extractive. But his proofs are not sufficiently conclusive. He has shown, however, that nutgalls contain acetic acid, and that the different gallates, when decomposed by sulphuric acid, emit the odour of acetic acid t.

Properties.

2. Gallic acid, when pure, is in the form of transparent plates or octahedrons. Its taste is acid, and somewhat astringent; and when heated it has a peculiar and rather unpleasant aromatic odour.

[·] Phil. Mag. xxiii. 74.

⁺ Jour. de Pbys. lxi. 117.

¹ Aun. de Chim. lx. 156.

3. It is soluble in 11 parts of boiling water, and in Chap. III. 22 parts of cold water. When this solution is heated, Solubility. the acid undergoes a very speedy decomposition. Alsohol dissolves one-fourth of its weight of this acid at the temperature of the atmosphere. When boiling hot, it dissolves a quantity equal to its own weight. It is oluble also in ether.

4. When exposed to the action of heat, it is sublimed, Action of but its properties are somewhat altered, as Bouillon La Grange has shown. In like manner the acid sublimed by Deyeux's process differs in its properties from the crystallized acid of Scheele and Richter. Deycux announced, that when the gallic acid is distilled it yields oxygen gas. When Berthollet repeated the experiment be obtained only carbonic acid. Bouillon La Grange has shown, that besides the carbonic acid, there comes over likewise a portion of heavy inflammable air, and that water is formed. By repeated distillations the whole acid may be decomposed and converted into these products. Hence it is obvious that gallic acid, like most of the other combustible acids is composed of oxygen, hydrogen, and carbon. The proportion of carbon, if we judge from the quantity of carbonic acid cvolved, must be very considerable.

5. Gallic acid in crystals is not altered by exposure Action at to the air. Neither oxygen gas, the simple combustibles, nor azote, seem to have any particular action on Its action on the metals has not been examined. When the solution of this acid in water is exposed to the air, it gradually acquires a brown colour, and the acid is destroyed; the surface of the liquid becoming covered with mouldiness.

6. It combines with alkaline bodies, separating the

carbonic acid if they were in the state of carbonates. The compounds formed have received the name of gallater; but hitherto have scarcely been examined.

Action on

7. In alkaline solutions it occasions no deposite, but when dropt into barytes water, strontian water, or line water, it gives them a bluish red colour, and occasions a flaky precipitate, composed of the acid combined with the earths.

Gallic acid occasions a precipitate when poured into solutions of glucina, yetria, and zirconia in acids. This property distinguishes these three bodies from all the other earths, none of which are precipitated from their solutions in seids by gallie acid †.

On metals.

8. Upon the metallic solutions it acts with considerable energy, changing the colour, and producing precipitates in many of them. Hence it is frequently used as a reagent to detect the presence of metallic bodies; but the difficulty of freeing it sufficiently from tannarenders it scarcely safe to trust the experiments hitherto made on that point. Richter has shown, that it is not capable of taking iron from sulphuric acid, as has been hitherto supposed, unless it be assisted by the action of some other body which has an affinity for sulphuric acid. He has endeavoured to show, too, contrary to the experiments of Proust, that it strikes a black with all the oxides of iron. Berthollet has more lately employed his ingenuity to establish the same doctrines,

It is not unlikely that these precipitates are occasioned by the prosence of a little tannin, and that they would disappear if the acid with pure.

If we except their solutions in carbonic and. Alumina is prospimed from acids by infusion of nut-galls, but not immediately

it has by no means silenced Proust, who has publish. Chap. III. experiments apparently decisive.

When it precipitates metallic oxides, gallic acid apears to act by bringing them nearer to the state of me-Ls, and some of them, as gold, are completely reduced.

SECT. IV.

OF TANNIN.

NUTGALLS contain several other ingredients besides History. allic acid; but one of the most remarkable and important is the substance called tannin, which will occupy our attention in this Section.

The first attempt at a regular examination of the proerties of nutgalls was made by Dr Lewis, during a et of experiments undertaken to ascertain the best mode f making ink *. He detected in them a substance rhich precipitates black + with the oxides of iron, and pagulates with isinglass ‡; but chemistry in his time ad not made sufficient progress to enable him either > separate or examine this substance. Deyeux was erhaps the first chemist who ascertained the peculiar ature of tannin. He pointed it out in his analysis of jutgalls as a peculiar resinous substance, but without ssigning it any name f. Seguin soon after engaged n a set of experiments on the art of tanning leather ||;

^{*} Phil sophical Commerce of the Arts, p. 377.

[†] Ibid. p. 346.

[‡] Ibid. p. 387.

[§] Ann. de Chim. XVII. 23.

[|] Ibid. xx. 38.

during which he discovered that tannis has the property of precipitating glue from its solutions in water, and a combining with the skins of animals. This led him a suppose it the essential constituent of the liquids employed for the purpose of tanning leather. Hence the names tannin and tanning principle given it by the French chemists; but it is to Mr Proust that we are indebted for the first investigation of the nature and properties of tannin, and of the methods of obtaining it in a separate state. Much curious and important information has likewise been obtained by the experiments of Mr Davy on the constituent parts of astringent vegetables, and on their operation in tanning. Fiedler, Richter §, and Merat Guillot ||, have also published interesting experiments on this difficult subject.

An elaborate dissertation on tannin was published by Trommsdorf in the summer of 1804 ¶; in which he analysed the opinions of Proust, and examined the properties of tannin with his usual industry. Many of his conclusions had been anticipated by Davy, whose hours, however, they serve to corroborate and confirm; but the recent labours of Mr Hatchett have formed a new era in the history of tannin. This sagacious philosopher, during a set of experiments on resins and bitumens, discovered a method of forming tannin artificially from almost every animal and vegetable body, and thus furnished chemists with the means of procuring it with facility in a state of purity. His disserta-

^{*} Ann. de Chim. xxv. 225.—xxxv. 32,—and xlii. 89.

[†] Pbil. Trans. 18:3, p. 233. and Jour. of the Royal Instit. vol. ii.

¹ Jour. de Chim. i. 86.

[§] Ibid. iii. 307. and 334.

[!] Ann. de Chim. xli. 323.

S Gehlen's Jour. iii. 111.

ne on an artificial tanning substance were read to the Chap. III. wal Society in the summer of 1805.

These discoveries will make it proper for us to die this subject into two parts. We are now in posstott of two kinds of tannin; one kind formed in ints by the processes of vegetation; another formed micually by the methods pointed out by Mr Hatchett: th of these appears capable of assuming different mofications, either from slight alterations in the constitus, or from the combination of small portions of foan bodies.

NATURAL TANNIN.

1. TANNIN exists in a great number of vegetable submees; but it may be procured most readily and in the matest purity from nutgalls and catechu.

NUTGALLS are excrescences formed on the leaves of Preparaoak by the puncture of an insect which deposits its as on them. The best are known by the name of Reppo galls, imported in large quantities into this untry for the use of the dyers, calico-printers, &c. they are hard like wood, round, often nodulated on the rface, of a bluish colour, and an excessively disagreele taste. They are in a great measure soluble in wawhat remains behind is tasteless, and possesses the operties of the fibre of wood. A very great proporon of water is necessary to carry off every thing soable. Deyeux found, that a French pound of putga'ls required set French posts of water, plant offerint partions one after the or t, and allowed to mace-Vol. 11.

rutgalla

standard, gives us about 150 English pints to a pound troy of nutgalls. But Trommsdorf exhausted the soluble part of nutgalls, by means of 40 times their weight of water, applied in three successive portions, each toutinuing two days on the galls, at the temperature of 60°r.

From the analyses of Deyeux and Davy, it follows, that the soluble part of nutgalls consists chiefly of fine ingredients; namely, tannin, extractive, mucilagels, gallic acid, and gallate of lime. Mr Davy found that 500 grains of Aleppo galls formed with water a solution, which yielded by slow evaporation 185 grains of matter. This matter he found composed of

130 tannin

31 gallic acid and extract

12 mucilage and extract

12 lime and saline matter

185 \$

So that the tannin constitutes rather more than two-third of the whole.

Prepara-

2. No fewer than five methods have been proposed to separate tan from the infusion of nutgalls, and procure it in a state of purity; but none of them asswers the purpose completely.

First. When a solution of muriate of tin is dropped into the infusion of nutgalls, a copious yellow precipitate immediately falls; which, when separated by fil-

Ann. de Chim. xvii. 12.

⁺ Gehlen's Jour. in. 113.

[‡] Extractive and mucilage are vegetable substances, which will be treated of in a subsequent part of this Work.

[§] Phil. Trans. 1803, 251

Chap. H.

pation and dried, assumes the appearance of a buff-cobured light powder. According to Mr Proust, who erst examined this powder, it is a compound of oxide of tin and tannin. If it be mixed with water, and a surrent of sulphureted hydrogen gas passed through it, alphuret of tin is formed, which remains insoluble; and the tannin, as it is separated from the oxide, discolves in the water. This water, when freed from the sulphuret by filtration, and evaporated to dryness, aves a brown coloured substance, which Proust condered at first as pure tannin . But as the infusion of satgalls contains a portion of extract, which is likewise precipitated by muriate of tin, it is obvious, that by this process we do not obtain pure tannin, but a combination of tannin and extract. Neither is the whole of the tannin precipitated; a portion of it combined with the exide remaining in solution, unless thrown down by an Mkalit. Mr Davy has rendered it probable that this recipitate contains also muriatic acid t. From these facts it is obvious that pure tannin is not obtained by his process.

Second, When the infusion of nutgalls, somewhat concentrated by evaporation, is mixed with a saturated solution of carbonate of potash, a yellows h white mater precipitates abundantly in the form of flakes. When his precipitate is dried, it assumes the form of a whitish owder, which was first examined by Deyeux, to thom we are indebted for the process f. Proust considers this precipitate as pure tannin; and accordingly

^{*} Ann. de Chim. 224, 226. \$ Phil. Trans. 1803, p. 249.

[†] Proust, Ann. & Chim. xli), Eg. 6 Ann. de C in . 18 . 19.

quid, which gave a black colour to oxysulphate of on, though it formed no precipitate with gelatine; it perefore contained gallic acid. Trommsdorf's experiments show us likewise that this method does not held pure tannin f.

Fourth. If time-water be mixed with an infusion of agalls, a copious precipitate falls. When this precipitate is treated with diluted nitric or muriatic acid, an fervescence takes place, the liquid becomes deep cobured; and when filtrated leaves behind it a substance f a brilliant black colour, which Merat-Guillot, to hom we are indebted for this process, considers as are tanning. But Mr Davy has shown that it must also ontain the extractive matter, which is thrown down in ombination with lime as well as the tanning. The fervescence indicates sufficiently the presence of cartonic acid; a proof that the precipitate is even still nore complicated. This method, then, is scarcely pre-trable to the former.

Fifth. The process practised by Trommsdorf, though not unexceptionable, appears to yield tannin in a state of greater purity than any other hitherto thought of. It as follows: Three parts of nutgalls were reduced to powder, and digested with 40 parts of water for three days, at the temperature of 66°, the mixture being frequently stirred. The whole was now passed through all en strainer, the liquid set apart, and the powdered outgalls remaining on the strainer were treated as before with 40 parts of water. This digestion with fresh

[@] Phil. 'ran. 503 140.

dan. de Gbim. zh. 323.

⁺ Gehlen's Jour. in. 127.

⁴ Phil. Tram. 1803. 262.

water was continued full four different infusions in all had been drawn off the nutgalls. The last of these was colourless, and produced no change upon the solution of iron. These infusions were mixed together, and evaporated jently down to one fourth in a porcelain basin. The liquid being now inuddy was passed through a dude lives cloth, by means of which a quantity of extractor was separated. The whole was then evaporated to the consistence of a jelly, and placed upon a flat porceian dish near a stove till it became quite dry. The brown coloured substance thus obtained was digested with thrice its weight of pure alcohol "; and this digestion was repeated in all three times, till the last portion of alcohol was found to contain no trace of gallic and. la order to make sure of removing the gallic acid, the dry residue was digested twice successively with alcohol containing ten per cent. of water. It was now considered as tannin in a considerable degree pure, but still contaminated with some extractive and mucilaginous matter. To get rid of these, the whole was dissolved in distilled water, and repeatedly evaporated to drynen, in hopes of rendering the extractive moseluble; but nothing was separated by this process. When the solution was left for some time in a warm place, a mould collected on the surface, which was removed, and ascribed to a portion of mucilage which had been present. The solution being now filtered, and evaporated to dryness, left a residue consisting of tannin in a state of considerable purity, but still contaminated with a quantity of sulphate of lime. To get rid of this salt, Mr Tromms-

Alcohol of the specific gravity 0.796 is at present considered as purpor free from water.

Chap 'II.

employed the following method: The tannin was polved in water, and carbonate of potash dropt in as as any precipitate fell. The liquid was separated m this precipitate by filtration, and mixed with acehe of lead. A powder fell, consisting of the oxide lead combined with tannin. This powder was washand dried; and being mixed with water, a current of aphureted hydrogen was passed through. By this cans the lead was separated and remained in combimion with sulphur, while the tannin dissolved in the tater. The liquid being now filtered, boiled, and evabrated to dryness, left a residue, which may be consibred as tannin in a state of as great purity as it can be socured from nutgalls *.

3. CATECHU, or terra japonica as it is also called, is From catesubstance obtained by decoction and evaporation from species of mimosa which abounds in India. It has reddish brown colour, an astringent taste, leaving an impression of sweetness; it is not altered by exposure the zir. There are two varieties of it; one from Bommy, which has the lightest colour, and a specific graviof 1.39; and one from Bengal, which is of the coour of chocolate; its specific gravity is 1.28 +. This abstance was examined by Davy, and found to consist hiefly of tannin combined with a peculiar species of structive. If the darkest parts of the catechu be seected, and infused in cold distilled water for a short time, the infusion, when evaporated to dryness, consists I tannin combined with a very minute quantity of extract. It may therefore be employed to ascertain the properties of tannin.

Gehlen, ili. 173, and 124.

book II.
Divis on II.
Properties

4, Tannin procured from nutgalls by Trommsdorf's method, is a substance of a brown colour, brittle, and breaking with a resinous fracture. Its taste is bitter and very astringent, like that of nutgalls.

It dissolves readily in water, both hot and cold, and forms a solution of a brown colour, which, from Trommsdorf's experiments, does not seem liable to become mouldy, nor to undergo a spontaneous decomposition when exposed to the air in a moderate heat.

Insoluble in aicohol.

Pure alcohol does not dissolve tannin; but it is readily soluble in alcohol diluted with water, even though the potential of water be but small. Thus alcohol, of the specific gravity 0.818, dissolves it, though it contains according to Lowitz's table, only to have been observed by Richter, and to have furnished him with the method for merly described, of procuring gallic acid in a state of purity.

Action of uzygen.

- 5. From the experiments of Proust, Davy, and Deyeux, we learn that it is capable of combining with oxygen, but at the same time it is either decomposed altogether, or its nature completely altered. Thus nitric acid converts it into a yellowish brown matter soluble in alcohol, and similar in its properties to an extract. Oxymuriatic acid produces similar effects; and Mr Proust has observed, that the peroxide of tin changes it also into an extract, perhaps by communicating oxygen.
- 6. The action of the simple combustibles on tannin has not been examined.

^{*} Davy, P il. Irans. 1203, 241.

7. The action of the metals upon tannin does not Chap. III. neem to be great; but almost all the metallic oxides have an affinity for it, and are capable of combining with it, the compound is usually nearly insoluble in water. Hence the reason why the infusion of nutgalls precipitates metallic solutions so readily. These compounds have been hitherto in a great measure overlooked by chemists. The following observations contain the facts at present known.

Action of metalac

When the peroxide of un or zinc is boiled in the infusion of galls, it acquires a dull yellow colour, and abstracts all the constituents from the infusion, leaving behind only pure water. The oxides thus combined with tannin, &cc. are partly soluble in muriatic acid. and the solution indicates the presence of tannin and gallic acid . When peroxide of tin is allowed to act upon the cold infusion, it abstracts all its constituents in a few days, but Mr Proust + affirms, that in that case the gallic acid is mostly destroyed, and a portion of the tannin brought to the state of extractive.

When the metallic salts are mixed with the infusion of galls, the precipitate consists of the metallic oxide combined with the tannin, the extract, and the acid of the infusion; and, according to Davy, it contains also a portion of the acid of the metallic salt 1.

Tannin does not seem to produce any change upon the solution of sulphate of iron: but when it is mixed with a solution of the oxysulphate of iron, a deep blue coloured precipitate immediately appears, consisting of

Davy, Phil Trans. 1803, 244. 1 Post. Iraas. 1803, 248.

⁺ Ann. de Chim. xlii. 92.

when dried, assumes a black colour. It is decomposed by acids.

When too great a proportion of oxysulphate of iron is poured into a solution of tannin, the sulphuric and, set at liberty by the combination of the iron and tannin, is sufficient to re-dissolve the precipitate as it appears; but the precipitate may easily be obtained by cautiously saturating this excess of acid with potash. When the experiment is performed in this manner, all the oxysulphate of iron which remains in the solution undecomposed is converted into sulphate. Mr Proust supposes that this change is produced by the tannin absorbing oxygen from the iron.

Tanner combines with gela-

8. One of the most important properties of tannin is the insoluble compound which it forms with glue or gelatine, as this substance is termed by chemists. It is therefore employed to detect the presence of gelatine in animal fluids; and, on the other hand, solutions of gelatine are employed to detect the presence of tannin in vegetable fluids, and to ascertain its quantity. Now although the compound of gelatine and tannin is insoluble in water, it is soluble both in the solution of tannin and of gelatine when sufficiently diluted. It is necessary, therefore, that the solution of gelatine, used to detect tannin, should be as concentrated as is consistent with its perfect fluidity; for glue, when gelatinous, does not act upon tannin. It is necessary also that it should be employed quite fresh; for when in a state of puttefaction, it loses its property of precipitating tannin.

[•] I find that even the addition of as nouth alcohol as is consistent with the gelatine remaining in solution, does not preserve it in a proper with for use.

Ir Davy has ascertained, that the best proportion for Chap. III. is a solution of 120 grains of isinglass * in 20 ounes of water. Care must be taken not to add an excess If the solution to the liquid from which the tannin is be separated; because the compound of tannin and relatine is re-dissolved by the solution of gelatine. Acparding to the analysis of Mr Davy, this compound, then dried in the temperature of 150°, is composed of

54 gelatine

46 tannin

100+

1. Potash and soda combine with tannin, and form Action of with it a compound less soluble in water than pure tanin, and which does not precipitate glue till the alkali saturated with an acid 1. Ammonia produces the time effects. The fixed alkalies occasion a precipitate a concentrated solutions of tannin, but ammonia throws Lown nothing .

When potash or soda is added to the infusion of nutpails, the liquid assumes a reddish-brown colour, and oses the property of precipitating gelatine till the alkali saturated with an acid. When the alkalized infusion is evaporated to dryness, an olive coloured mass remains, of a faint alkaline taste, which deliquesces in the ir. Ammonia produces the same effect upon the infusion of galls; but when the mixture is exposed to the

^{*}Isinglass is glue or gelatine nearly pure, as has been shown by Mr.

⁺ Pb. Trans. 1803. p. 235, and 250.

¹ Trommsdorf, Gehlen, in 144.

Of carths.

heat of boiling water, part of the ammonia flies off, a precipitate falls, consisting of most of the tannin and gallic acid, while the extract remains in solution §.

10. When barytes or lime water is poured into a solution of tannin, a precipitate falls, consisting of tannin combined with the earth, and the solution becomes nearly colourless. The precipitate dissolves with difficulty in water, and does not act upon the infusion of gluetil the earth is saturated with an acid.

When newly precipitated magnesia is agitated with the infusion of tannie, it unites with a portion of it, and forms a smoky brown powder insoluble in water, but soluble in acids. Alumina produces the same effect, and forms a similar compound †.

When barytes, strontian, or lime water, is poured into the infusion of gails, an olive coloured precipitate falls, which consists not only of the tannin, but also of the extract, and most of the gallic acid combined with the earth. When magnesia is mixed or boiled with this infusion, it combines with all its constituents; the gallate remains mostly in solution, and gives the liquid a green colour; while the tannin and the extract form with the magnesia an insoluble compound, and give ita dirty yellow colour. Alumina in small quantity produces exactly the same effect; but when used in a greater proportion, it separates all the constituents of the infusion ‡.

When the earthy carbonates are boiled in the infusion of galls, they separate the tannin and the extract, while

[§] Davy, Phil. Irins 1802, n. 241.

Tronumsderf, Gehlen' J r. i.i. 145. † Ibid.

[‡] Davy, Pbil. 1rans. 1803, p. 241.

bey combine with the acid, and form with it a salt hich remains in the liquid, and gives it a green co-

Chap. III.

Of acida.

11. Most of the acids have the property of combiing with tannin, and of forming solutions more or less huble. Acetic, phosphoric, oxalic, and malic acids, ccasion no precipitate when dropt into a concentrated olution of nutgalls. Arsenic acid produces a copious recipitate, soluble in boiling water, and precipitating due after the acid has been neutralized by an alkali. Turistic acid likewise produces a precipitate mostsoluble in hot water. The same remark applies sulphuric acid. But this acid alters and gradually secomposes tannin. Nitrie acid produces no precipite in the infusion of nutgalls. The mixture becomes tot, and assumes a red colour, which gradually chanles to a yellow. By the action of this acid a bittersted substance is formed, which possesses the properles of malic acid +.

12. When tannin is distilled, it yields an acid liquor, which blackens solutions of iron, because it contains a little tannin unaltered: there comes over also some empyreumatic oil, and a voluminous coal remains beaind, amounting to 1/2 of the tannin distilled 1.

13. Such are the properties of the tannin of nutgalls, far as they have been ascertained. The difficulty of procuring it in a state of purity renders some of them

[•] Davy, Flul. Trans 1803, p. 241.

[†] Trommsdorf, Gehlen's Jour. ni. 142.

[†] Proust, Ann. de Chim. Exev 35.

Book II. Division 11.

Writing

ambiguous, and has induced chemists to employ a reagent the entire solution of nutgalls.

This solution is employed in considerable quantums, by the diers, and it forms the principal ingredient a writing ink. It is not known at what period thus apportant liquid came into use, but the ink of the accients was composed of very different ingredients, as ing analogous to the ink used by the printers at pasent. We are indebted to Dr Lewis for a valuable at of experiments on the best mode of making ink.

This liquid consists of a solution of sulphate of me in the infusion of nutgalls, and seems to owe in blick colour chiefly to a combination of the tannin with he exide of iron, or perhaps with the sulphate, and proje also to the combination of gallic acid and oxide of us The follest black is produced when equal weight of green vitriol and galls are used; but the ink very som fades. To make it permanent, the galls ought to be thrice the weight of the vitriol. No other solution of iron but the sulphate forms with nutgalls a full black. When the mixture of the infusion of nutgalls and great vitriol is diluted with much water, the black manu precipitates, and forms a sediment not again soluble. The addition of log wood increases the blackness of the The following formula was ascertained by Dr. Lewis to yield the best ink.

Logwood..... ounce

Nutgalls in powder....3
Green vitriol1

Water to 2 quarts

Boil the logwood and nutgalls in the water, adding new liquid in proportion to the evaporation, then strain through a cloth, and add the vitriol to the water, addang at the same time from one to two ounces of gum Chap. III. erabic. As soon as these have dissolved, the ink is fit for use . Some recommend the addition of a little doves in powder to prevent mouldiness.

14. Mr Proust has announced it as his opinion, that there exist various species of tannin in the vegetable kingdom, differing from each other like the oils, resins, &c. He has even enumerated several of these variepies, and pointed out their characteristics +.

It is by no means unlikely that this opinion is well Species of Sounded, and the experiments of Mr Hatchett serve to confirm it: But it is impossible to admit it as demonstrated till a process be discovered for obtaining tannin in a state of purity; for the differences between the varieties pointed out by Proust may be owing to the presence of foreign substances which disguise its properties. At any rate, this subject will come under our consideration more properly in the Second Part of this Work, when we enumerate the different vegetable substances that contain tannin.

II. ARTIFICIAL TANNIN.

THE important discovery, that a substance possessing Discovery. similar properties to the tannin of nutgalls may be formed artificially by a very simple process, was made by Mr Hatchett in the course of a set of experiments on the slow carbonization of vegetable bodies, and detailed by him in two papers read to the Royal Society in 1805.

Book II. Division II. Formation.

1. To form this artificial tannin, we have only to be gest diluted natric acid on charcoal till the whole, it nearly the whole, is dissolved. Mr Hatchett usually employed 100 grams of charcoal, and 500 gram & nitrie acid of the specific gravity 1:40, diluted with tage its weight of water. On heating this mixture man and mateass a considerable effervescence was produced, 24 much nurous gas esca, ed. After two days digest a more nitric acid was added, and the digestion was tinued till the solution was complete. The scheme thus obtained was transparent, and of a dark brown colour; which being evapolated to dryness, leaves a brown coloured mass. This is the artificial tannan. To the it from the last portions of nitric soid. Mr Hate of found the best method was to dissolve it repeatedly awater, and evaporate cautiously to dryness with a gene tle heat.

By this method 100 grains of charcoal were converted into 120 grains of artificial tannin; but of these.

Mr Hatchett supposes three grains to be moisture.

Properties.

2. Tannin thus prepared is a substance of a brown colour, has considerable lustre, and breaks with a sitteeous fracture. Its taste is bitter and highly astrongent. It has no smell.

It dissolves readily in cold water, forming a true parent solution of a deep brown colour. Alcohol ala dissolves it *.

The alcohol used by Mr Hatchett was probably weak. Then tion of pure alcohol has not been tried. Were it solvable in it, this was form a marked distinction between natural and artificial tann. I at the effect of alcohol of she specific gravity 0.800, the ptrongest 1 had my possession. It formed a pale yellow solution.

3. The solution of artificial tannin immediately precitates glue, or isinglass, from water. The precipitates ces are more or less brown according to the strength the solutions, and always insoluble both in hot and ld water. To ascertain the proportion in which these ries unite, Mr Hatchett dissolved 50 grains of artifitannin in 4 02, of water, and then precipitated by cans of a solution of isinglass. Forty-six grains of be tannin were precipitated in combination with eightyne grains of isinglass. Hence the precipitate is comessed of about

36 tannin 64 isinglass

The solution containing the four grains not precipitated y the isinglass being evaporated to dryness, left a light Tittle substance of a pale brown colour, smelling strong. of prepared oak bark, especially when dissolved in ater. The solution tasted bitter, and from its properies approached nearer to the vegetable matter called extractive than tannin; a proof that even the artificial annin prepared from charcoal is not quite free from foreign bodies.

Action of

4. When sulphuric acid is added to a solution of ar- acids. tificial tannin, a copious brown precipitate falls, soluble in boiling water, and capable of throwing down gelatine. Muriatic acid produces precisely the same effect. Artificial tannin dissolves readily in nitric acid, but is not altered in its properties, though that acid be repentedly distilled off it. In this respect it differs very materially from all the species of natural tannin hitherto examined, which were found by Mr Hatchett to be en-

tirely decomposed by nitric acid, though with different degrees of facility.

Alkalier.

both fixed and volatile. When dissolved in amount, evaporated to dryness, and again dissolved in water, to new solution does not precipitate gelatine unless the previously mixed with a small portion of muriate and. A proof that it still retains the ammonia in combination. When the fixed alkalies are added to a solution of artificial tannin, the colour is immediately deeper, and after some hours the solution becomes turbid. Cubonate of potash produces the same effect, and the some time a brown magma is deposited.

Earths,

6. The alkaline earths unite with artificial tanta, and form compounds little soluble in water. Hence a forms a precipitate when mixed with nitrates of line, barytes, &c.

Oniden.

7. It precipitates likewise most of the metallic exide from their solutions in acids. The colour of the precipitate is usually brown, inclining to chocolate.

Action of

8. When artificial tannin is thrown upon a hot irm, it emits an odour similar to that of burning feathers. When exposed to a graduated heat in a retort, there passes over, in the first place, a portion of water; and that is succeeded by a little nitric acid, from which it is difficult to free it completely. A little yellow liquor next makes its appearance; and upon raising the fire, amnoniacal gas is disengaged with 'great rapidity. This is followed by the evolution of carbonic acid gas, together with a small portion of gas, which seems to possess the properties of azote. A bulky coal remains in the retort, amounting in weight to 0:425 of the original to-

in. This coal being burned, left some brown ashes, Chap, III. chiefly of lime.

From this decomposition by the action of fire, it is Constitubyious, that artificial tannin is composed of oxygen, azote, hydrogen, and carbon. The last ingredient obriously predominates; though the proportions have not hitherto been ascertained.

Such are the properties of the artificial tannin from barcoal, as far as they have been hitherto ascertained by Mr Hatchett, to whom we are indebted for all the acts above detailed. From the experiments of this indefatigable chemist, we learn that every kind of chargoal yields it equally, from whatever substance it has been formed, whether vegetable, animal, or mineral, provided it be in the state of charcoal. But the action of nitric acid on charcoal, though the readiest and easiest process, and that which yields the greatest quantity, is not the only one by means of which artificial tannin may be formed. Mr Hatchett has pointed out two others, by means of which certain vegetable substances may be converted into tannin.

9. The first of these consists in digesting nitric acid Second vawith certain substances that appear to contain an uncommon portion of carbon in their composition; indigo, for instance, and many of the bodies called resins. Indigo dissolves readily in diluted nitric acid, and the solution, when gently evaporated to dryness, leaves an orange-coloured mass of an intensely bitter taste, soluble in water, and possessing the property of forming an insoluble precipitate with gelatine. Hence it resembles tannin; but it acts more feebly on the metallic salts than tannin from charcoal. A similar substance was procured from common resin by digesting it for a long

time in diluted nitric seid. Most of the resins and goal resins gave a similar product; but no tannta could be produced from the gums.

Third va-

10. The other process for procuring artificial tanial consists in dissolving resins and camphor in sulphure acid, digesting the solution till it becomes black, and then precipitating by throwing it into cold water. A black powder falls. If this powder be digested in all cohol, a brown substance is taken up, which is soluble both in water and alcohol, forms an insoluble precipitate with gelatine, but acts only feebly on oxy-sulphare of iron. Camphor by this process yields nearly half in weight of a brown resinous-like matter, which possesses the property of forming an insoluble precipitate with gelatine; and when digested with a little nitric acid, becomes precisely similar to tannin from charcoal.

Thus there are three species of artificial tannin. 1. Tannin procured by the action of nitric acid on this coal. 2. Tannin, by digesting nitric acid on indige and resins. 3. Tannin, by dissolving resins or camphoria sulphuric acid *.

Such are the properties of the colorific acids.

act with most energy on metallic solutions, forming precipitates which vary in colour according to the metal. It is this property which renders them of so much importance in a chemical point of view. The colour of the precipitates which each of these bodies forms with the different metallic bodies, as far as is known at pre-

sent, may be seen from the following TABLE.

Chemical uses of the coloratic acids.

^{*} See Mr Flatchett's papers, Phil. Truns. (805 and 1806. From which all the facts respecting artificial tannun have been taken.

Chap. Hi.

fron	O. or sulphur	Black		Blue	Blue	Mack	Brown
Cupper,	Black	Black		Green-	Вгочи	C) B comes	Olive
Constants.						Blue	
Indum,				colos rl. 4		O Nec mea	
Rhodism.		0		0			
Mercury. Palladium, Rhodium,	Dark	Black ?		Ol.ve			
Mercury.	Black	Black	C	White	Orange Jellow	Orange	Yellow
Silver.	Black	Black	White	White	Yel brown	Yel brown	Yellow
Platinum.	Reduced	Black	0	0	0	0	
Cold.	Reduced	Black	Reduced	Yell	Reduced	Reduced	Reduced
	Sulphureted Reduced Reduced Black	Hydrcsulph. Black of potash.	Prussic acid. Reduced	Prusviate of potasb.	Gallic acid. Reduced	Infusion of Reduced	Tannin from Reduced

[TABLE continued.]

	Nickel	Tin.	Lead.	Zinc.	Bismuth. Antimony Tellurium.	Antimony	Tellurium.	Arsenic.	Cobalt.	Manga- nese.
Sulphureted hydrogen.	0	Brown	Black	White	Black	Orange		Yellow	0.	0
Hydrosulph. of potash.	Black	Black	Black	White	Black	Orange	Brown	Yellow	Black	White
Prussic acid.								:		·
Prussiate of potash.	Green	White	White	White	White	0	0	White	Brown yellow	Yellow white
Gallic acid.	White	0	White	0	Orange	White		0	 O	0
Infusion of Galls.	Grey	Brown	White	0	Orange	White	Yellow	0	Yellow.	
Tannin from Charcoal.		Black	Brown	0		Yellow		Yellow?		

Chap.	III:

	Chromium.	Molybde- num.	Molybde- Urmium.	Tungsten.	Titanium.	Columbi- um.	Niccola- num.	Cesium.
Sulphureted hydrogen.		Choco-			0			· :0
Hydrosulph. of Potash.	Green	Reddish brown	Brown yellow		Bottle green	Red cho-		Brown
Prussicacid.		·	•			• •		
Prussiate of Potash.	Green		Brown	Brown	Grass	Olive	•	White
Gallic acid.							•	0
Infusion of Galls.	Brown		Choce-	Straw	Reddish brown	Orange		0
Tannin from Charcoal.							,	

Book II. Biv.sion II.

CHAP. IV.

OF COMPOUND COMBUSTIBLES.

THE dompound combustibles are almost all competed of carbon and hydrogen, or of carbon, hydrogen, and They are a very numerous class of bodie, comprehending the greater number of animal and vegetable substances, and of the products obtained from these substances. But the present state of chemistry does not permit us to take the term in that comprehensive sense: The investigation of vegetable and animal bodies is too incomplete; their properties are too imperfectly known to allow us to introduce them into the first principles of the science; and the utility of the greater number of them as chemical instruments is too inconsiderable to warrant any such introduction, even if their properties were completely investigated. For these reasons, it will be proper to treat in this Chapter of those compound combustibles only which are employed in chemistry as instruments of investigation, reserving the remainder for the Second Part of this Work, These may be reduced under five classes; namely,

Number of compound combustibics.

- 1. Alcohol
- 4. Fixed oils
- 2. Ether
- 5. Bitumens
- 3. Volatile oils

The properties of these bodies form the subject of the fellowing Sections.

SECT. I.

OF ALCOHOL.

HE liquid called alcohol, or spirit of wine, is obtained History, by distilling wine, beer, and similar fermented liquors. These liquors appear to have been known in the ear-Hest ages. The Scripture informs us that Noah planta vineyard and drank wine; and the heathen wrias are unanimous in ascribing the invention of this liwor to their earliest kings and heroes. Beer, too, seems have been discovered at a very remote period. It was in common use in Egypt during the time of Heronotus". Tacitus informs us that it was the drink of the Germans t. Whetler the ancients had any method procuring ardent spirits from these or any other limors does not appear. The Greeks and Romans seem have been ignorant of ardent spirits altogether, at least we can discover no traces of any such liquor in meir writings. But among the northern nations of Europe, intoxicating liquors were in use from the earliest wes. Whether these liquors resembled the beer of the Germans we do not know.

At what period these liquors were first subjected to distillation is unknown; though it can scarcely have preceded the time of the alchymists. The process is simple. Nothing more is absolutely necessary than

⁺ De Morib. Germ. ch. xxiu. # Lib. il. p. 77.

over is ardent spirits. It is certain, at least, that the method of procuring ardent spirits by distillation was known in the dark ages; and it is more than probable that it was practised in the north of Europe much earlier. They are mentioned expressly by Thaddæus, Villanovanus, and Lully *.

It is by the distillation of fermented liquors that ardent spirits are obtained; and they receive various names according to the nature of the substance employed. Thus brandy is procured from wine, rum from the fermented juice of the sugar cane, which and gin from the fermented infusion of malt or grain. Now ardent spirits, whatever be their name, consist almost entirely of three ingredients; namely, water, pure spirit or alcohol, and a little oil or resin, to which they owe their flavour and colour.

Rectified spirits.

1. When these spiritous liquors are redistilled, the first portion that comes over is a fine light transparent liquid, known in commerce by the name of rectified spirits, and commonly sold under the denomination of alcohol or spirit of wine. It is not, however, as strong as possible, still containing a considerable portion of water.

Preparation of alcohol.

The method usually practised to get rid of this water is to mix the spirits with a quantity of very dry and warm salt of tartar †. This salt has a strong attraction for water, and the greatest part of it is insoluble in alcohol. It accordingly combines with the water of

[·] Bergman, iv. art. ii. 4.

[†] Impure petzeh not fully saturated with carbonic acid.

the spirit; and the solution thus formed sinks to the Chap. IV. bottom of the vessel, and the alcohol, which is lighter, swims over it, and may easily be decanted off; or, what is perhaps better, the solution of potash may be drawn off from below it by means of a stop-cock placed at the bottom of the vessel . The alcohol, thus obtained, contains a little pure potash dissolved, which may be separated by distilling it in a water bath with a very small heat. The spirit passes over, and leaves the potash behind. It is proper not to distil to dryness. This process is first mentioned by Lully. The liquid procured by means of it has been usually distinguished by the name of alcohol.

Alcohol is said to have been discovered, or at least first accurately described, by Arnold de Villa Nova, who was born about the end of the thirteenth century. This chemist, who was professor of medicine at Montpelier, first formed tinctures, and introduced them into medicine t.

The specific gravity of spirits, as highly rectified as Strength of possible by repeated distillations, seems to be about common al-0.820, at the temperature of 60°; but the alcohol of commerce, which is nothing else than rectified spirits, is seldom under '8371. By means of salt of tartar Muschenbroeck brought it as low as .815; but, in general, the alcohol concentrated by that process is not under 821, owing to the weakness of the spirits employed. Even at the specific gravity '815 the alcohol is by no

[.] See this process described by Hoffman as new, in his Observationes Phys -Chym Select. p. 36. published in 1732.

He has said also to have been the first who obtained the oil of turpentine. He procured it by distilling turpentine, and employed it as a solvent of resina.

Book 15. Diesson II. means pure, still containing a considerable portion of water. Dr Black, by repeated distillations off mutate of lime, obtained it as low as '800; but it was Low's of Petersburgh who first bit upon a method of absolute purity, or at least very nearly so. His process was published in 1790°, and the same year Richter made known another, by when the same purification was accomplished with still greater facility †.

Method of procuring pure alcoto-

The process of Lowitz is as follows: Take a quantity of fixed alkali t perfectly dry, and still warm, and newly fill with it a retort. Upon this pour such a quantity of alcohol (previously brought to 821 by means of salt of tartar) as can be absorbed by the alkali completely, so that the whole shall have the appearance of a solid mass without any alcohol swimming above. In general, the proportion ought to be two parts alkali and one part alcohol. Allow this mixture to remain for 24 hours, and then distil by a heat so moderate, that about two seconds elapse between the falling of the drops of alcol of frem the beak of the receiver. When this interval mercases, the receiver mus be changed; for it is a sign that all the strong alcohol has come over. What comes over next is weaker. By this process Lowitz obtained alcohol of the specific gravity 0:701 at the temperature of 68°.

When Richter repeated the experiment of Lowitz, he reduced the alcohol to the specific gravity 0.792 at the temperature of 68°, but could not bring it any lower.

Crell's Annais, 1796, i. 195.

^{1 1}bid. ti. \$22.

[#] I presume in the state of a b-carlonate

He found, upon trial, that the following method, which is much more expeditious, answered equally well with that of Lowitz: He exposed a quantity of the salt called muriate of lime to a red heat, reduced it to powder, and introduced it while yet warm into a retort, and poured over it at intervals a quantity of alcohol, of 0.821, nearly equal to it in weight. A violent heat was produced. The retort was put upon a sand-bath, a receiver adjusted, and the liquid made to boil. The salt was dissolved, and formed with the alcohol a thick solution. The portion that had passed over into the receiver was now poured back, and the whole distilled by a gentle boiling nearly to dryness. The alcohol that came over was of the specific gravity 0.792 at the temperature of 68°.

2. Alcohol obtained by these processes is a transpa. Properties. rent liquor, colourless as water, of a pleasant smell, and a strong penetrating agreeable taste. When swallowed it produces intoxication. Its properties differ somewhat according to its strength. When procured by Lowitz's or Richter's process, we may distinguish it by the name of pure alcohol, as no method known can deprive it of any more water. When the specific gravity is higher, the alcohol is contaminated with water, and the proportion of that liquid present increases with the specific gravity. Chemists, in general, have employed this impure alcohol, or this mixture of alcohol and water, in their experiments; and as they have too often neglected to point out the specific gravity of the spirit used, we are still in some measure ignorant of the properties of this important liquid .

[·] Fahrenheit was one of the first who ascertained some of the re-

Book If. Division II. Action of heat and cold. 3. Alcohol is exceedingly fluid, and has never been frozen, though it has been exposed to a cold so great that the thermometer stood at --69°. Indeed, Mr Walker sunk a spirit of wine thermometer to --91°, without any appearance of congelation.

It is a very volatile liquid. Fahrenheit found that alcohol of the specific gravity of about '820, at the temperature of 00°, boiled when heated to 170°°. When of the specific gravity '800 it boils at 1731°. At this heat it assumes the form of an elastic fluid, capsus of resisting the pressure of the atmosphere, but which condenses again into alcohol when that temperature is reduced. In a vacuum it boils at 50°, and exhibit the same phenomena: so that were it not for the pressure of the atmosphere, alcohol would always exist a the form of an elastic fluid, as transparent and invisible as common air. This subject was first examined with attention by Mr Lavoisier†. The fact, however, had been known long before.

Combination with water, 4. Alcohol has a strong affinity for water, and is miscible with it in every proportion. The specific gravity varies according to the proportion of the two h-

markable properties of alcohol with exactness. His alcohol was of the specific gravity 825 at the temperature of 48°. See Phil. Trans. 124 vol. 1220. Four croy informs us that the specific gravity of the most highly received alcohol is 0.9213, without specifying the temperature. The extremity of Baume's hydrometer for spirits (according to Nicholson's table, Quarto lour, 1. 19) answers to the specific gravity 817, temp 55°. This may be considered as beyond the strength of the alcohol used. In Germany, before Lowitz's experiments, the wronger alcohol seems sell out to have exceeded 0.521 at 68°; and in the country of its commonly considerably weaker. The highest point of Clark's hydrometer corresponds with alcohol of about 834 at 30°. In Lews scatter the purest accohol of the specific gravity 820— Newson's Cham.

* Ped. Iran. 1222. vol axim p 8. . . | Jose. de Phys. 178:

ids combined; but, as happens in almost all commations, the specific gravity is always greater than mean of the two liquids; consequently there is a intual penetration: and as this penetration or condention varies also with the proportions, it is evident at the specific gravity of different mixtures of alcohol water can only be ascertained by experiment. As spiritous liquors of commerce are merely mixtures salcohol and water in different proportions, and as their sength can only be ascertained with precision by means of their specific gravity, it becomes a point of great importance to determine with precision the reportion of alcohol contained in a spirit of a given pecific gravity: and as the specific gravity varies with temperature it is necessary to make an allowance or that likewise.

The importance of knowing with precision the proextion of alcohol contained in spirits of every specific ravity, has induced many different persons to make speriments in order to ascertain this point with exactpes; but as they set out from alcohol of very different grengths, it is not easy to compare their results with seh other. We have seen that the pure alcohol, by owitz's process, is of the specific gravity .791 at the imperature of 68°. That chemist mixed various proportions of this alcohol with given weights of pure waer, and after adowing the mixtures to remain for 24 ours, took the specific gravity of each at the temperaure of 68°. The following Table exhibits the result these experiments. The first two columns contain e proportion of alcohol and water (in weight) mixed ogether, and the third the specific gravity of the mixare at 68°. I have added a fourth column, containing

Chap. IV.

Book II. the specific gravity at 60°, the temperature commonly preferred in this country *.

Table of the strength of alcohol of parious denlities.

		C					
100	Parte.	Sp. G	ravity.	100 I	Mt.	Sp. C	sary.
Alooh.	Wat	at '680.	at 60°.	Algob.	-Wat.	at 68°.	at 60%
100	1	791	796	70	50	868	871
90	1	794	798	Gġ	31	810	874
28	2	797.	801	·08	32	872	875
97	3	800	804	67	33	875	879
96	4	803	807	66	34	877	880
95	5	805	809	65	35	880	883
94	Ø	808	812	64	30	882	886
93	7	811	815	63	37	885	889
92	8.,	813	817	62	38	887	891
91	9	816	820	61	39	889	893
90	10	818	822	60	40	892	896
89	11	821	825	59	41	894	898
88	12	823	827	58	42	896	900
87	13	820	,830	57	43	899	903
86	14	828	832	56	44	901	904
85	15	831	835	55	45	903	906
84	16	834	838	54	46	905	908
83	17	836	840	53	47	907	910
82	18	839	. 843	52	48	909	912
81	19	842	.846	51	49	912	915
80	20	844	848	50	50	914	917
79	21	847	851	49	51	917	920
78	22	849	853	48	52	919	922
77	23	851	855	47	53	921	924
76	24	853	857	46	54	923	926
75	. 25	856	860	45	55	925	928
74	26	859	803	44	56	927	930
73	27	861	865	43	57	930	933
72	28	863	867	42	58	932	935
71	29	866	870	41	59	934	937

^{*} Crell's Annels, 1796, i. 202.

d		_						
î	190 E	AFTE.	Sp. Gi	ravity.	100	Parts	Sp. G	envity.
ľ	Alcoh.	Wat.	at 680.	at 60°.	Akob	Wat.	at 68°.	at 60°.
ı								
1	40	50	936	0.7 0	19	81	974	975
ł		61	938	941	18	82	976	
ı	38	62	940	943	17	83	977	
ı	37	63	942	945	16	84	978	
1	36	B B	944	947	15	88	880	
ı	35	65	946	949	266	86	981	
l	10.0%	66	948	951	13	87	DEC.	
l	33	67	950	953	12	88	965	i
1	32		952	955	11	89	986	
ľ	31	69	954	957	10	90	987	
1	30	70	956	958	9	91	988	
l	29	71	957	960	8	92	8.00	- 1
I	28	72	959	962	7	93	991	1
ı	27	73	961	963	6	94	992	
I	26	74	963	965		95	994	
I	25	75	965	967	4	96	200	
I	24	76	966	968	3	201	997	
1	23	77	968	970	2	98		
	22	丁芳	970	972	1	99	999	
l	21	197.52	971	973		100	1000	
	20	80	973	974				

The importance of this object, both for the purposes revenue and commerce, induced the British Governent to employ Sir Charles Blagden to institute a very minute and accurate series of experiments. An account these was published by Blagden in the Philosophi-I Transactions for 1790; and a set of Tables, exhi- Gipin'staiting the result of them, was drawn up by Mr Gilpin, bles. ho had performed the experiments, and published in he Philosophical Transactions for 1794. The follow-Table, extracted from these, contains the specific ravity of different mixtures of alcohol and water at Vol. II. Dd

water. From the preceding Table, we see of .825 is composed of .89 pure alcohol
11 water

Chris	LIV.
-	لخصضم

water.	helt roo alcoholt ou afe droi roo alea re 15 water. 10 water. 25 water	30 water.	35 water.	40 water	45 water
84995 85957 84769 85729	86625 87357 88059 86587 87357 88059	88701	89244	9.003 t	90345
8550	m -	88491	89013	80017	90127
84076 85042	8738	_	88026	84114	89684
84502	1999	87796	883,43	6824	89458
83362 84334 8	5193 65976	-	8:6:8	840	0 0
124 84092 8	36	-	81105	88254	In I
83851	85490 8		81466	88018	85538
82396 83371 8	84248 85036 85757	86411	106	87590	22
82150 83126 8	84001 84797 85518	86172	86787	87360	87889
81900 82877 8.	3153 84550 85272	œ 	86542	87114	87454
7 82 39 8		00000	98 0	50870	37421

COMPOUND COMPUTIBLES.

Book II. Division II.

Real Specific Gravities at the different Temperatures.

sleebol water.	16686.	08180	93592	93382	11116	02963	92158	8546	08333	11120	1881	91189	11510	00610	0000
hel 100								<u> </u>		-	O 01		-	_	4
co alcohe yo water.	14186.	93541	9334	93131	01676	10126	09770	02283	09060	91840	91098	91465	9184	02010	90805
alcoholico	41486.	93274	93078	92859	92647	98180	92825	92 010.	20710	91560	01340	91186	9606	14100	28500
roo alcohol 80 water.	10186	93626	92783	92570	92158	98145	91083	91715	91403	01810	\$1010	90886	#990#	90443	51800
too alcohol 75 water.	68836.	92680	98110	9536	15086	91637	91688	91400	19110	90059	90183	90558	90349	90119	9999
	.98563	98955	15186	18610	91723	91503	51887	99016	L+806	2008	90385	60806	8000	80708	20834
too alcohol 65 water.	11256.	92009	91109	91584	91370	9116	78000	70706	90484	90252	12006	80843	11068	89190	80158
too alcohul	.91847	91640	91 +28	11216	16606	90708	90549	90328	90106	89872	89639	80460	80830	89003	88769
too alcoho: SS wa er.	91440	14216	91026	90812	9059d	90367	90144	89920	89695	89464	89225	80043	8881-7	88588	1888
So water. SS wa er. 60 water. 65 water. 70	.91023	90-11	90506	90380	09106	89933	89707	89479	89252	81068	88781	88605	88376	88146	37015
Heat	300	35	0+	. 45	50	55	90	65	20	75	80	•5	8	95	100

Chaps IV.

S alcohol 90 po water 110	alcehol	Heat, 1:00 sie holl 95 akehol 90 alerhol 85 alerhol 80 alerhol 100 water, 1:00 water, 1:00 water.		175 alcohol 170 alcohol 100 water.	70 alcohol 100 water.	of alcohol	60 alcohol	55 alcohal fob water.
94441 -94615 -94920	.8492	10	-95173	*95429	-95681	-95944	.96 209	.96410
94249 94484 94734	9473	4	94688	95246	95502	95772	84096	90-15
94058 94295 94547	9454	1	94802	95060	95.128	95602	95879	96159
93860 94096 94348	943	8+	94005	17840	95143	95423	95705	95993
93658 93897 94149	941	40	94414	94683	91958	95243	95534	95831
93452 93696 939	939	3948	94213	94486	94767	95057	D-857	95662
93247 93493 93749	937	40	94018	94296	94579	04810	18156	95493
91040 93285 935	935	940	9 1822	04000	98816	94689	95000	95318
92828 93076 9339	933	[- [-	93616	93898	94193	94000	94814	95139
92613 92565 931	931	64 05	93413	93695	686.6	94301	94623	94957
02393 9.646 0291	029	111	93×01	93488	99169	04102	04431	94768
_							I	
				_				_

Real Specific Gravities at the different Temperatures.

Heat	50 alcohol 100 water.	45 alcohol 10 : water.	io alcol·ui	35 a cohel 100 wat r	30 aic .!. 100 w 1		ico warer, 100 water. I O were 100 water 100 water	ts alcoh l	to richol e alcohol too water. 100 water.	c alcohol
300	•96719	L9696.	-97200	.97418	.97635	.97860	80186	.98412	+0886-	.99334
35	96579	96840	91085	97319	97556	10816	9801B	98397	68 804	99344
40	96434	90196	19696	97220	97472	91137	98033	98373	98195	99345
45	96280	96563	04896	97110.	1384	99916	08616	08338	98774	99338
50	96156	96420	80196	96905	97284	91580	97920	1.8.93	98745	99316
55	95966	96272	96575	11896	07181	97500	07847	98249	2018 0	99284
00	05804	96122	96437	96752	97074	974:0	וררדפ	98176	98654	99244
65	95635	59056	9:1288	96620	06090	60816	92926	98106	98:54	99194
10	95469	95802	96143	96484	96836	97203	97596	86.38	98527	18166
75	95292	95638	95987	14896	8u196 .	92026	81405	97043	98454	99066
80	1 95111	95467	95826	96195	96508	89096	91385	21845	98367	98991

By the excise laws at present existing in this coun- Chap. IV. the duty on spirits is levied by the bulk, estimated particular strength, to which all spirits of whatever egth are brought by means of a set of tables calcud. This strength is regulated by a very inconvenihydrometer, invented in 1730 by Mr Clarke . which time it has undergone various changes and provements. This instrument was adapted to a very hard kind of language, which had previously come into expon use among dealers, and seems from them to me made its way into the excise laws. A mixture of bulks of alcohol and water was called proof spiand sometimes double spirit. Clarke's hydrometer paists of a thin co, per ball, terminating above in a slender stem, and below, in a metallic button, to p ic perpendicular. It is so light as to swim in pure bohol. There is a mark upon the midd'e of the stem; a small weights accompany the instrument, to be put in order to suit the various temperatures of the li-There is a weight marked proof, so adjusted et when it is placed upon the stem along with the toper weight for the temperature, the stem sinks till mark on the middle of it is on a level with the surce of the liquid.

From an act of parliament passed in 1762, we learn at, at the temperature of 60°, the specific gravity of coof spirit ought to be 0.916. But Clarke's hydroneter, loaded with the proper weights, sinks to the ark indicating proof in spirits of the specific gravity 920, at the same temperature. From the table of

See the first description of it in Phil. Trans. 1710, v. l. xxxv . p. 277.

Lowitz, given above, we learn that a mixture of equal weights of water and pure alcohol has the specific gravity '917 at 60°. The legal proof spirit differs but it. the from this; Clarke's proof consists of 49 part of pure alcohol and 51 of water. The near coincidence of these numbers to the truth, indicates a considerable approach to accuracy. Had Sir Charles Blagden made choice of an alcohol of '800 for his standard, the specific gravity of proof spirit would have been found to deviate very little from that of Clarke. But if by proof spirit were to be understood equal bulks of water and alcohol, as was doubtless the case, then both the legal and Clarke's proof differ very far from the truth.

The strength of spirits stronger than proof, or overproof, as it is termed, is indicated on Clarke's hydrometer by the bulk of water necessary to reduce a given bulk of spirits to the specific gravity denominated proof. Thus, if one gallon of water be required to bring 20 gallons of the spirits to proof, it is said to be one to 20 over proof; if one gallon of water be necessary to bring 15, 10, 5, or 2 gallons of the spirits to proof, it il said to be one to 15, one to 10, one to 5, one to 2 respectively, over proof, and so on.

The strength of spirits weaker than proof, or ander proof, is estimated by the quantity of water which it would be necessary to abstract, in order to bring the spirits in question up to proof. Thus, if from 20 gallons of the spirits one gallon of water must be abstracted to bring it to proof, it is said to be one in 20 under proof; if from 15, 10, 5, 2, &c. gallons of the spirits, one gallon of water must be abstracted to bring it to proof, it is said to be respectively one in 15, one in 10, one in 5, one in 2, &c. under proof.

Degrees of Clarke's by-. drometer.

The following Table points out the specific gravity Chap. IV.

pirits of the various strengths indicated by Clarke's rometer at the temperature of 60° .

Degrees.	Sp. gravity.	
1 in 2	9644	
1 - 3	9549	
1 — 4	9458	
1 — 5	9424	
1 — 6	9385 ·	
1 - 7	9364	
1 — 8	9344	
1 — 9	9334 . `	
1-10	9320	
1 15	9280	
1 20	9265	
Proof	9200	
1 to 20	9162	
i — 15	9135	
1 10	9107	
1 - 9	9093	
1 - 8	9071	
1 — 7	9047	
1 - 6	9006	
1 5	8961	
1 — 4	8913	
1 — 3	8817	
1 — 2	8590	
Alcohol	8338	

remove all confusion from the duties on spirits, by ought to be levied by the weight, and not by the lk. This might be done with sufficient accuracy

Wilson's Tables, p. 1 .

either by actual weighing, or by constructing tables indicating the weight from the bulk and specific gravity.

Action of

5. Neither common air nor oxygen gas has any acgen on alcohol in moderate temperatures, whether in the liquid or gaseous state; but in high temperatures the case is different. When set on fire in the open air it burns all away with a blue flame, without leaving Boerhaave observed, that when the vaany residuum. pour which escapes during this combustion is collected in proper vessels, it is found to consist of nothing but water. Junker had made the same remark: and Dr Black suspected from his own observations, that the quantity of water obtained, if properly collected, excceded the weight of the alcohol consumed. servation was confirmed by Lavoisier; who found that the water produced during the combustion of alcohol exceeded the alcohol consumed by about \$th part . Mr Saussure junior has shown that 100 parts of alcohol when burnt, yield 132 parts of water †. A proof that it contains a considerable proportion of hydrogen as a constituent.

> When the vapour of alcohol is mixed with oxygen gas in the proper proportion, the mixture detonates when presented to a lighted taper, or when fired by electricity, as a mixture of oxygen and hydrogen gases This fact seems to have been first observed by Dr Ingenhousz, or at least his experiments on ether appear to have led to it ‡. The density of the vapour

Par. 1781, p. 493.

on's J. mil, axi. 263.

Love his Exterior in et Chiero etinei sen divers Object de Plante 12:

alcohol is considerable, and hence the quantity of Chap. IV. kygen necessary to consume it is great. The proacts of the combustion are water and carbonic acid .

6. Alcohol has but little action on the simple comastibles. On hydrogen, carbon, and charcoal, it does of appear to have any effect.

It dissolves a little phosphorus when assisted by heat. Phosphathis phosphorized alcohol exhales the odour of phosdureted hydrogen gas. When a little of it is dropt to a glass of water, a flame instantly makes its apearance, and waves beautifully on the surface of the ater +. This phenomenon, which is occasioned by he emission of a little phosphureted hydrogen gas, m only be observed when the experiment is performed a dark room.

When sulphur and alcohol are brought into contact Sulphurethe state of vapour, they combine and form a reddish bloured liquid, which exhales the odour of sulphureted Tydrogen. This compound was first formed by the Count de Lauraguais, who employed the following proess. Some flowers of sulphur were put into a large class cucurbite, having a glass vessel in its centre conmining alcohol. A head was adjusted, the cucurbite placed in a sand-bath, and heat applied. The sulphur was volatilized, and the acohol converted into vapour at once. These meeting together in the head, united and formed the red liquor wanted t. It was supposed by

cd alcohol.

Cruikshanks, Nicholani's Quarto Jour. v. 205.

⁺ Brugnatelis, Ann. de Clim 221v. 71. This experiment was first made Boyle. See Shaw's Boyle, wi. 196,

^{\$} Mew. Par. 1758 p 9.

chemists that sulphur cannot be dissolved in alcohol, except by a similar process, but from the late experiments of Favre, this does not appear to be the case. He digested, during 12 hours, one part of towers of sulphur in eight parts of alcohol, of the specific gravity 0.837, in a heat not sufficient to produce boding. The alcohol assumed a yellow colour, and acquired the smell and taste of sulphureted hydrogen. Another partion of the same alcohol was digested for a month, cold, on sulphur. The effect was the same. On trying alcohol of various strengths from .817 to .867, he found, that the alcohol acted with more energy in proportion to its strength.

The sulphureted alcohol prepared by Laursquair method, contains about to the oulphur. The sulphur is precipitated by w. ter.

Action on alkahes,

7. Alcohol dissolves the fixed alkalies very readily, and forms with them a reddish-coloured actid solution. It is from this solution only that these alkalies can be obtained in a state of purity. When heat is applied to it the alcohol may be distilled over. It appears however, to be partly decomposed; but the nature of the products has not been accurately ascertained. Ammonia also combines with alcohol with the assistance of heat: but at a temperature somewhat below the boiling point of alcohol, the ammonia flies off in the state of gas, carrying with it, however, a little alcohol in selution.

Earths.

8. None of the earths are acted upon by alcohol, un-

^{*} See the Elonous de Chamie of the Dijon Academy, iii. 287. Fourere 't Connoissances Chymiques, viel. † Gehlen's Jour. ii. 343.

s strontian and barytes be excepted. It absorbs aout its own weight of nitrous gas, which cannot afterwards be expelled by heat 1.

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9. Of the acids, the sulphuric, nitric, and oxymuri- Acids, tic, decompose alcohol; but all the other acids are soable in it, except the metallic acids, phosphoric acid, and perhaps also prussic acid.

10. Alcohol is capable of dissolving a great many sales. wline bodies. A considerable number of these, with be quantities soluble, is exhibited in the following Tables .

I. Substances dissolved in large Quantities.

Names of the Substances.	Tempera-	240 parts of Al- cohol dissolve		
Oxysulphate of iron Nitrate of cobalt †	54·5°	240 parts		
copper †	54.5	240 240 ' 300		
magnesia † Muriate of zinc †	180°5 54°5	694 240		
alumina † magnesia †	54·5 180·5	1313 240		
copper †	180·5 154·5	240 240		
Nitrate of zinc decomposed †				
iron decomposed † bismuth decomposed †				

[?] Prientley, i. 379.

[#] Morveau, Jour de Phys. 1785.

[§] Withering, Paul, Trans. luxii. 236.

II. Substances dissolved in small Quantities.

Names of the Sul-stances.	240 parts of Alcohol at the boiling tempera- ture dissilve		
Muriate of lime †	240 perts		
Nitrate of ammonia †	214		
Oxymuriate of mercury	212		
Succinic acid †	177		
Acetate of soda +	112		
Nitrate of silver +	100		
Refined sugar +	59		
Boracic acid †	48		
Nitrate of soda †	23		
Acetate of copper †	18		
Muriate of ammonia +	17		
Superarseniate of potash +	9		
Oxalate of potash †	7		
Nitrate of potash †	5		
Muriate of potash †	5		
Arseniate of soda +	4		
White oxide of arsenic +	3		
Tartrate of potash †	1		
Nitrate of lead \			
Carbonate of ammonia §			

III. Substances insoluble in Alcobol.

Sugar of milk	Sulphate of potash †
Borax+	soda †
Tartar †	magnesia†
Alum †	Sulphite of soda
Sulphate of ammonia †	Tartrate of soda and
lime †	potash
barytes §	Nitrate of mercury †
iron †	Muriate of lead †
copper †	silver ‡
silver †	Common salt +
mercury †	Carbonate of potash
. zinc †	soda

These experiments were made chiefly by Macquer and Wenzel*. The alcohol employed by Macquer was of the specific gravity 0.840. Wenzel does not give he density of his alcohol; but as he compares it with hat of Macquer, we may suppose it nearly of the same trength. As the solubility of salts depends upon the trength of the alcohol employed, the experiments of hese chemists must be considered as defective, because they have confined themselves to one particular density. This defect is in part supplied by the following very valuable Table of Mr Kirwan's, constructed from his own-experiments †.

^{*} Verwendtechaft, p. 300. The solubility of all the salts marked † was ascertained by Wenzel; those marked †, by Macquer; and those marked §, by Withering.

[†] On Mineral Waters, p. 274.

Solubility of Salts in 100 parts of Alcohol of diffuse

Salta.	Alsohel of				
	0.900	0.872	0.848	0-834	CF 1854
Snlpbate of soda	0,	0.	0.	0.	0.
Sulph, of magnesia	1.	1.	0.	0.	0-
Nitrate of potash	2'76	1		0.	0
Nitrate of soda	10.2	6,			0.
Muriate of potash	4.62	1.66		0.38	0.
Muriate of soda	5.8	3.67		0.2	_
Muriate of ammon.	6.5	4.12		1.2	
Muriate of magne-	21.25		23.75	36.25	50.
Muriate of barytes	1.		0.29	0-185	C-09
Ditto crystallized	1.26		0143	0.35	0.06
Acetate of lime	2.4		4.12	4.15	4.88

Its flame tinged by When alcohol containing certain saline bodies in solution is set on fire, its flame is often tinged of different colours according to the body. Thus nitrate of strontian tinges it purple; boracic acid and cupreous salts tinge it green; muriate of lime gives it a red colour; nitre and oxymuriate of mercury a yellow colour.



11. Different opinions were entertained by chemists out the nature of alcohol. Stail thought that it was Composiimposed of a very light oil, united by means of an acid a quantity of water. According to Junker, it was imposed of phlogiston, combined with water by means In acid. Cartheuser, on the other hand, affirmed that contained no acid, and that it was nothing else than are phlogiston and water. But these hypotheses were ere assertions supported by no proof whatever. Las sier was the first who attempted to analyse it.

He set fire to a quantity of alcohol in close vessels means of the following apparatus: BCDE (fig. 13.) wessel of marble filled with mercury. A is a strong was vessel placed over it, filled with common air, and pable of containing about 15 pints (French). Into wessel is put the lamp R filled with alcohol, the hight of which has been exactly determined. On the sek of the lamp is put a small particle of phosphorus. he mercury is drawn up by suction to the height IH. his glass communicates by means of the pipe LK with other glass vessel S filled with oxygen gas, and placed per a vessel of water T. This communication may be at up at pleasure by means of the stop-cock M.

Things being thus disposed, a crooked red hot iron are is thrust up through the mercury, and made to such the phosphorus. This instantly kindles the wick, id the alcohol burns. As soon as the flame begins to yow dim, the stop-cock is turned, and a communicaon opened between the vessels S and A; a quantity oxygen gas rushes in, and restores the brightness of he flame. By repeating this occasionally, the alcohol my he kept burning for some time. It goes out, how-

Sent 1

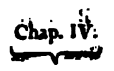
= last, notwithstanding the admission of cay

After the combustion, there was found in the remel 115:41 cubic inches of exchanic acid gas weight of which was 78:1102 grains troy. There likewise found a considerable quantity of water it resert, but it was not possible to collect and weight [Lavoisier, however, estimated its weight at aggrains; as he concluded, with reason, that the of the substances employed were still in the way when the whole contents of the vessel consisted of beauc acid gas and water; therefore the carbonib gas and water together must be equal to the axign and alcohol which had been consumed.

But 78°1192 grains of carbonic acid gas contained acid gas contained on the Lavoisier's calculation. 55°2°9 (at oxygen grains, however, of oxygen grains must have each oved in forming water.

water, 01038 grains of hydrogen gas require, in order to water, 01038 grains of hydrogen gas; and the quarter formed by this combination is 411265 grains of water found 89 005 grains of water figure vessel; therefore 47193 grains of water must record ready formed in the alcohol.

[.] Men. Por. 1781.



6.03 hydrogen 47.83 water

76.7*

Such were the consequences which Mr Lavoisier drew From his analysis. He acknowledged, however, that There were two sources of uncertainty, which rendered This conclusions not altogether to be depended upon. The first was, that he had no method of determining the quantity of alcohol consumed, except by the difference of weight in the lamp before and after combustion; and that therefore a quantity might have evaporated without combustion, which, however, would be taken into the sum of the alcohol consumed. But this error could not have been great; for if a considerable quantity of alcohol had existed in the state of vapouf in the vessel, an explosion would certainly have taken place. The other source of error was, that the quantity of water was not known by actual weight, but by calculation.

To this we may add, that Mr Lavoisier was not warranted to conclude from his experiment, that the water found in the vessel, which had not been formed by the oxygen gas used, had existed in the alcohol in the state of water: he was intitled to conclude from his data, that the ingredients of that water existed in the alcohol before combustion; but not that they were actually

^{*} Mem. Par 1781.

combined in the state of water, because that combine tion might have taken place, and in all probability at partly take place, during the combustion.

34 carbon17 hydrogen

100

A result certainly far from the truth. From the experiments of Cruikshanks, made by detonating a mixture of the vapour of alcohol and oxygen gas, it follows, that in alcohol the proportion of carbon is to that of bydrogen as nine to one. But this is not sufficient to give us the component parts of alcohol with precision.

The subject has been lately resumed by Saussure junior, who has published a very elaborate set of experiments on the analysis of alcohol. The alcohol which he employed was obtained by Richter's process, and, of course, of the specific gravity '792, at the temperature of 65°. He employed three different methods of analysis; the first method was similar to that of Lavoisier. He burnt a quantity of alcohol in common air mixed with oxygen gas, and ascertained the weight of alcohol and exygen consumed. The only sensible products were carbonic acid and water, the first of which he

[.] Nichaise's Jures!, v. 205.

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carbon 36.890 hydrogen 15.814

100.000+

He found, that when a considerable quantity of alcoholows burnt, and the water formed collected, it contained portion of ammonia. Hence he concluded that azote one of the constituents of alcohol.

His second method of analysis was to mix the vapour of alcohol with oxygen gas when the thermometer stood about 70°, and to detonate the mixture by means of electricity. He estimated the quantity of alcoholic vapour present by means of Mr Dalton's formula, explained in another part of this Work. To make the mixture capable of detonating, he was obliged to add a small quantity of hydrogen gas. The result of the experiment was, that 18.19 grains of alcohol in the state of vapour consumed 75.88 cubic inches of oxygen gas. There were formed 46.69 cubic inches of carbonic acid gas, together with a portion of water. From these data the calculated the component parts of alcohol as follows:

The weights and measures mentioned in these experiments are French.

¹ Nicholson's Jour. XII. 225.

Hork II. Division 1f. oxygen 37.36 carbon 46.82 hydrogen 15.82

This result differs from the preceding, and ought! more accurate. It deviates, however, very far from conclusions drawn by Cruikshanks from a similar of experiments.

The third method of analysis was to decomposite alcohol by passing its vapour through a red hot lain tube: the products were a little charcoal, oil, partly in crystals, partly fluid, a portion of holding in solution acetic acid, ammonia, and a acid which resembled benzoic, and a great quar heavy inflammable air, to which Saussure ga name of exycarbureted hydrogen gas, and which his analysis, he considered as a quadruple composition, carbon, hydrogen, and azote. The folis the composition of alcohol as deduced from the

These results, though probably as exact as the j state of our knowledge will enable us to go, are be considered as exact. The absolute alcohol of

Nicholson's Jour. axi. 259.

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coubtless contains a portion of water; and our ignoof the quantity renders every attempt to analyse of uncertain. I do not consider the analysis of the arbureted hydrogen as made by Saussure to be ex-The proportion of azote which he infers was proan error in the experiment. The presence of pp is, however, in the water from alcohol, seems to ne oof that azote actually exists in alcohol, though reportion of it is so small that it may have been from some foreign body accidentally present. at it contains oxygen, has been proved by a very set of experiments performed by Messrs Four-Vauquelin. When equal parts of alcohol I phuric acid are mixed together, the sulphusuffers no change; but the alcohol is decomtheing partly converted into water and partly er. Now it is evident that the alcohol could not converted into water unless it had contained

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OF ETHER.

alcohol is mixed with sulphuric acid and sether acids, and the action of the acid is assisted the alcohol is decomposed, and converted partly

[·] Michalson's Jour. i. 391.

Book II. Devision II Varities of other. into a very light volatile fragrant liquor, known by the name of ether. The properties of the ether obtained are supposed to vary a little according to the acid casployed: accordingly every particular kind is duraguished by the acid used in its preparation. The the ether obtained by means of sulphuric acid, while sulphuric ether; that by means of nitric acid, while ether. It will be proper to consider each of these species separately,

I. Sulphunic Ethen.

History.

The method of making sulphuric ether is described in the dispensatory of Valerius Cordus, published a Nuremberg about the year 1540; from which Courd Gesner transcribed it into his Theraurus Euonymide Pamediis Secretis, published in 1552, where it is calcul Oleum Vitrioli dulce*. It appears to have been known,

^{*} Whoever will consider the formula given for preparing this 2000 Fittriols dulie by Gesner, will be satisfied that it was very different less the duterfied seeds of the moderns, and that it must have been a minimum of elsabel, ether, and rever out of wine. The following is the passage of free ner, as quoted by Hoffman, from whom has been taken the hinertal facts respecting the knowledge of other possessed by the alchymatical weiters. " Recipe vint ardenin acervini de ter mbil mati uneral quaque, plei vitrioli austeri tantundem, misce in veneti - i vitro, & pone fin zurbitam parvam angusto utificio, & luto uptituo e tificium claute, de mitte ita per integriim mensem aut dunt. Deinde effunde in queut. tam, our sit immediate annexum alradicum, citjus figuram sabjasemas Pone deinde in paream fornacem, ac dinudiam tjus partem cinere corte postea applica recipientem & luto juncturam claude diligenter, & extrahpurcian sex vini profentis quas infudinti. Ut vero tutius hae fiat, pere n balpeum Marin, sie sohim vinum ab que oleo ascender. L'um estratorie autem per balneum infunes uncies ter van mei, pane id, gnod rendung

though not in a state of purity, both to Basil Valen- Chap. IV. But in the writings of chemists cine and Paracelsus. published about the end of the 17th century, I have not been able to find any traces of it *, except in those of Mr Boyle. He was evidently acquainted with it, as appears from different passages of his writings +, though he no where describes it particularly. But it was a paper in the Philosophical Transactions for 1730, by a German who called himself Dr Frobenius, describing several of its most singular properties, that first drew the attention of chemists to this curious liquor 1. In this paper it first received the name of ether. The German chemists long distinguished it by the name of naphtha.

1. Sulphuric ether is usually prepared by the follow- Preparaing process !: A mixture of equal parts of alcohol and

🗫, in formecem, ut arena mediam cucurbitæ partem atringat, ac novo & vacuo recipiente coque nun magno applicato, luto juncturam dil genrer claude. Accende deside modestum igners, & censim extrahe omnem Lumiditatem que relicia est in cucurboa, di nec nibil humidi amplius in fur do appareat, adh b ta semper maxim cura & diligentia, ut ignem ita modereris, ne ebulliat usque ad alembici canalem. Nam si hune ebuilitin attigerit, sedare 1 on potes, 1 eque prob bere, quin in receptaculum egreenatur, ac totum oleum perdat; solet enim facillime chulure. Tum videbis duo conticere in eo, aquesin videbeet humorem ac pinguem; segregable vero unum ah altero statim, ita ut nih l aqueum in oleo reliaquatur, nam aqua illa oleum corrempir; segreg dum oleum usui reserva."

• The Gleum Vitrials suice of Lemery, for apstance, is very different from that described by Gesner. (See his Cours as Llyone, p. 502.)

+ See Shaw's Boyle, i. 530, and 1 269; where the process for making ether, and some of its must remarkable properties, are cetailed at length.

? Plot Trans. xxxvi. 283. This paper is little else than a rhapsody in the alchym stical style. At the end of it there is a note by Mr Godfrey [Hankwaz', Mr Boyle's operator, mentioning the experiments formerly made upon a Ly Mr Beyle and S.r Japae Newton.

* Frobenius' process was first published in the Philosophical Transactions " "

Bark II. Danision II sulphuric acid is put into the retort, to which a large receiver is then luted. It is proper to surround the is ceiver with ice, or at least with cold water. Healt applied; and as soon as the mixture boils, the energomes over and is condensed, and runs in large sits down the sides of the receiver. As soon as it amount to one half of the alcol of employed, the process it is be stept. The ether thus obtained is not quite purpalmost always containing a little sulphurous acid.

This acid may be separated by pouring the cust on a little potash, and distilling it over again by means of a moderate heat. Mr Dize officials that black oxide of manganese produces this effect still more completely than potash. All that is necessary is to max a quantity of mis black oxide to powder with the impure ether, and to let it remain for some time, agreeing it occasionally. The sulphurous acid is converted into sulphurous and combines with the marganese. The other is then to be distilled over by the heat of a water bath?.

The separation of the liquid from the sulphurous acid, with which it is mixed, is called the rectification of the other. The usual method, and I may add the best, is the following, first employed by Mr. Wolter Fill three-fourths of a bottle with the impute ether, add a little water and a portion of slacked lime. Against the bottle with violence, and keep it for some time in water before taking on the cirk. If the smell of the acid be not removed, add a little more lime, and against a second time. Decant off the ether into a retornal and distill it over †.

f Trous, des, de Cour. zim 256.

[·] Int. de l'hy. abet agil.

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The ether procured by this process is not quite pure, en after all the acid has been removed. ruon of liq aid that comes over during the distillation sneedy alcohol impregnated with a little ether. Comon ether is in reality a mixture of ether and alcohol. De usual method of separating this liquid is by mixing e ether with water, and then proceeding to distillation ath a very moderate heat. But Mr Lowitz has shown at this method does not succeed. The following proess yielded him an ether much purer than any that had een previously obtained. Into 16 parts of ether, of the secific gravity '775 in the temperature of 60°, he threw by powdered salt of tartar, till the last portions were o longer wetted by the liquor. The mixture being allowed to digest, the ether was then drawn off. Its speafic gravity was now only .7.6. By this means it was eprived of the water which it contained. To remove the alcohol, dry powdered muriate of lime was thrown, to the liquid in the same manner, as long as it would issolve. On standing, the mixture separated into two ortions; the alcohol holding the salt in solution sunk the bottom; the ether swam on the surface. When eparated from the inferior liquor, its specific gravity ras now only '632 in the temperature of 60°. It was erefore much purer than any former ether described chemists, since it never before had been procured ghter than 0.725 . The ether thus prepared conins a little of the salt, from which it may be freed by isullation. But in that case its specific gravity in-

[•] Lowitz, Crell'e Annale. 2796, i. 429.

Properties,

creases. The reason seems to be, that the purest portion of the other assumes the form of clastic fluid.

2. Ether thus obtained is a limpid and colourless to quor, of a very fragrant smell, and a hot pungent mue.

It is so volatile that it can scarcely be poured from one vessel to another without losing a considerable pertion of it by evaporation. When poured out in the open air, it disappears in an instant; and during its evaporation produces a very considerable degree of cold. If a glass vessel containing water, and surrounded with a cloth, be dipt into ether, two or three times, and the ether each time be allowed to evaporate from the cloth, the water in the glass freezes. In the open air ether boils at 95°, and in a vacuum at —20°. Were it not therefore for the pressure of the atmosphere it would always exist in the gaseous state.

When exposed to the open air it speedtly assumes the gaseous form. This happens, for instance, if a little of it be poured into a glass phial. The vapour of other displaces a considerable portion of the air of the phial, and is not soon disapated. Ingentious a har shown that the specific gravity of this vapour is very considerable.

Mr Dalton has found it 2.25, the specific gravity of common air being 1. According to the estimate of Saus are at the temperature of 72; , two French ounces of other, when converted into vapour, occupy the space of about a French cubic foot † If this estimate be corsect, 100 cubic inches of otherial vapour at that tem-

[†] Nicholson's Jour. 221, 335.



See his aveilles Auperionees, p. 180.

rature weigh only 45.15 grains troy, which would the its specific gravity only 1.45. This estimate is eatly below that of Mr Dalton, and in all probability low the truth.

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Ether, when exposed to a cold of—40°, freezes and ystallizes †.

3. Neither oxygen gas nor common air produce any fect upon ether in moderate temperatures; but in high imperatures the case is very different. Ether is exedingly inflammable, and when kindled in the state vapour burns with rapidity, with a fine white flame, id leaves behind it a trace of charcoal. During its imbustion carbonic acid is generated. How well sover it has been rectified, it always exhibits traces of alphuric acid ‡.

When ether is admitted to any gaseous body standing over mercury, it always doubles the bulk of the as, as Dr Priestley first observed. If oxygen gas, in expanded by ether, be presented to a lighted candle, we ether burns with great rapidity, but produces no uplosion. But if one part in bulk of this expanded tygen be mixed with three parts of pure oxygen gas, and kindled, a very loud explosion takes place: the oducts are water and 2½ parts of carbonic acid. Mr ruikshanks, to whom we are indebted for this instructure experiment, ascertained, that one part of the vapour ether takes 6.8 parts of oxygen gas to consume it empletely; and from the relative proportions of the

[†] Fourcroy and Vauquelin, Ann. de Chim. xxix. 289

¹ Scheele, 11. 108.

Cruikshanks, Nicholson's Journal, v. 205.

two products, he has shown that the carbon which the contains is to its hydrogen as five to one.

According to Dalton, one part of other by weight in quires for its combustion 3 parts of oxygen: the products are 1½ parts of water, and 2½ of carbonic acid; Saussure junior has lately endeavoured to ascertain the constituents of other, by mixing a known quantite is etherial vapour with oxygen gas, detonating the minute, and estimating the proportion of oxygen commend and of carbonic acid formed. The following is the result of his experiments. Ether is composed of

Composirice,

> carbon 55.2 hydrogen 22:14 oxygen 19:66

> > 100.00

What renders this result not to be implicitly depended on is, the probability that the other of Saussure, which was of the specific gravity 0.717, still contained a partion of alcohol. We see from it, however, that sulphosic other contains much less oxygen and much more earbon and hydrogen than alcohol.

Ingenhousz was the first who ascertained that the vapour of other detonates with common air and oxygen
gas. His account of the experiment was first published
in a letter to Dr Priestley, in one of the original volumes of that philosopher on Air, and likewise in the
ooth volume of the Ph losophical Transactions. His
method was exceedingly simple. A single drop of

¹ New System of Oranized Philosophy, 5, 195.

Nicholson's James, 221, 328

ether, let fall into a bottle holding about 10 cubic inches Chap. IV. of air, gives it the property of detonating. Too much ether destroys the detonation. With oxygen gas the same method succeeds .

When ether in the state of vapour is made to pass through a red hot porcelain tube, it is decomposed completely, and a great quantity of carbureted hydrogen gas is obtained †. Saussure junior has lately repeated the experiment with precision. He passed 1103 parts of the other through a red hot porcelain tube; the products were as follows: $5\frac{1}{4}$ parts of charcoal in the tube, 3 parts of volatile oil crystallized in thin scales and smelling of benzoin; 43 parts of a volatile oil nearly black, partly fluid and partly of the consistence of boney; 3 parts of water and 948 parts of heavy inflam-The loss amounting to 100.75 parts was mable air. chiefly owing to the escape of oil in the state of vapour ‡.

4. Ether does not combine with water in any proportion: when the liquids are shaken together, they separate again; but the water retains a portion of the ether, while the ether on the other hand remains united to a part of the water. From the experiments of the Count de Lauraguais, we learn that ten parts of water take up one of ether §. Alcohol, on the other hand, unites with ether in any proportion whatever.

^{*} See Ingenhousz' Experiences, p. 171.

[†] Dutch Chemists, Jour. de Pleys. xlv. 184.

² Nicholson's Journal, xxi. 323.

Mem. Par. 1758. From his experiments it seems to follow, that the portion taken up by water is not other, but a substance which may be obtained in crystals by evaporation.

only on phosphorus and sulphur.

Is is capable of dissolving a small proportion of phosphorus. The solution is transparent; but the addition of a little alcohol to it renders it milky. This furnishes us with a method of ascertaining whether either be sopphisticated with alcohol 1.

Ether was supposed incapable of acting on sulphus, except when both were in the state of vapour, according to the experiments of Lauraguans; but Favre has shown that a solution may be obtained by digestary flowers of sulphur in cold ether, and that the solvent power of the ether is promoted by exposure to the light. By a month's digestion, he dissolved nearly out part of sulphur in 12 of sulphuric other; the solution was nearly colourless, but had the taste and smell of sulphurited hydrogen †.

that have a weak affinity for oxygen when mixed with their solution in acids, as gold and silver. It dissolves the muriate of gold and the oxymuriate of mercury.

7. It is probable that it has no action on fixed alkalies and earths; but it combines, or at least mixes readily with ammonia.

It absorbs nitrous gas in considerable quantity.

8. Sulphuric acid seems capable of convening it into a peculiar kind of oil known by the name of sweet oil of wine.

If we fill a bottle capable of holding three or four



l Brugnatelli, Ann de Clem, xxiv. 73.

[†] Gehlen's Jame, 19, 227.

moglish pints with oxymuriatic acid gas, taking care to expel the water as completely as possible, and then throw into it about a dram, or half a dram, of good ther, covering its mouth immediately with a piece of light wood or paper, in a few seconds white vapour will be perceived moving circular in the bottle: this will be soon followed by an explosion accompanied with same; at the same time a very considerable quantity of charcoal will be deposited, and the bottle will be found to contain carbonic acid gas . The action of the other acids upon other has not been examined with attention.

9. Ether dissolves the fixed and volatile oils, bitumens, those at least which are fluid, and resins; but it does not act upon gum +.

10. Chemists entertained various opinions respecting Theories of the nature of ether. Macquer supposed that it was tion. merely alcohol deprived by the acid of all its water. But it was generally believed that the acid entered partly into its composition. Scheele published a set of experiments on ether in 17821; from which he drew as a consequence, that during the process the alcohol is deprived of phlogiston. These experiments were varied, and carried still farther by Pelletier; who adopted the theory of Scheele, modified according to the discoverics of Lavoisier. According to him, ether is alcohol combined with oxygen. This theory was embraced by the greater number of chemists; and it was supposed that the alcohol obtained the new dose of oxygen from

Craikshanks, Nicholson's Jour. v. 205.

[†] Elemene de Chymte by the Dijon Academy, iii. p. 327

¹ Schoele, il. 105.

been lately examined with much care by Fourtrey and Vanquelin. These ingenious chemists have concluded from their experiments, that during the process the moshol is completely decomposed, and that other is composed of the same ingedients as alcohol, but combined in different proportions. Either, according to them contains a greater proportion of hydrogen and oxygen and a smaller proportion of carbon than alcohol.

The theory of these chemists was disputed by Law det + and Dabit 1, who endeavoured to prove that orggen is always necessary for the formation of ether-This they did by repeating the experiments on other which had been formerly made by Scheele. They distilled a mixture of sulphuric acid, black oxide of manganese, and alcohol. The sulphuric acid was not decomposed, as in common cases, no charcoal was deposeted, no gas came over, the black oxide lost part of its oxygen, and the quantity of ether obtained was greater than usual. According to Dabit, ether contains a smaller proportion of hydrogen, and a greater proportion of oxygen and carbon, than alcohol. These objections were almost immediately answered by Fourcroy and Vauquelin, who proved that ether obtained by means of the black oxide of manganese possesses very different properties from sulphuric ether. Consequently is formation and composition cannot destroy their concusions respecting the formation and composition alphuric ether f. From the preceding analysis of Sam-

^{4 1}bid. 319.



Ann. de Clim 22is . 203, 2nd Nicholson's Jeer. i. 39 t.

[†] Bar. de Ciem, LERIV. 282.

^{\$} Ibid. 2.2217. 289.

mire it follows, that ether contains less oxygen that al- Chap. IV. cohol, but more carbon and hydrogen.

11. As the action of sulphuric acid on alcohol, and Action of the formation of ether, is one of the most interesting phenomena in chemistry, and as it may enable us to form more precise ideas, both respecting the composition of alcohol and ether, it will be proper to examine it with attention.

stilpharic acid on alco-

. When four parts of sulphuric acid and one part of 1. Olefland alcohol are mixed together, and a moderate heat applied, the mixture blackens, boils violently, and a great quantity of gas is disengaged. This gas, which was first examined by the Dutch chemists, received from them the name of olefiant gas. It has been described in a former part of this Work, under the name of supercarbureted kydrogen *.

What remains in the retort after the disengagement of this gas is chiefly sulphurous acid blackened with charcoal, and probably also some vegetable acid.

When equal parts of sulphuric acid and alcohol are 2. Ozalia mixed together, the phenomena which take place are considerably different. If the mixture be made cautiously, and allowed to remain at the common temperature for about 30 hours, orystals of oxalic acid often form in it +.

A combination of two parts of sulphuric acid and one of alcohol elevates the temperature to 201°, becomes immediately of a deep red colour, which changes to a black a few days afterwards, and emits a smell percepably ethereal.

Book II. Der sinn H. When a mixture of equal parts of alcohol and supplied acid is exposed to the action of heat in a proper apparatus, the following phenomena take place, as he been ascertained by Fourcroy and Vauquelin.

3. Ether.

When the temperature is elevated to 208°, the and boils, and emits a vapour which becomes condensed by cold into a colourless, light, and odorant liquor, which from its properties has received the name of ether. If the operation be properly conducted, no permanent gains disengaged until about half the alcohol has passed over in the form of ether. Until this period there puses absolutely nothing but ether and a small portion of water, without mixture of sulphurous or of carbone acid.

If the receiver be changed as soon as the sulphuron soid manifests itself, it is observed that no more ether is formed, but the sweet oil of wine, water, and accoms acid, without the disengagement hitherto of a single bubble of carbonic acid gas. When the sulphuric acid constitutes about four-lifths of the mass which remains in the retort, an inflammable gas is disengaged, which has the smell of ether, and burns with a white only flame. This is what the Dutch chemists have called chiftant gas. At this period the temperature of the fluid contained in the retort is elevated to 230° or 234°

When the sweet oil of wine ceases to flow, if the receiver be again changed, it is found that nothing more
passes but sulphurous acid, water, carbonic acid gas;
and that the residuum in the retort is a black mass, consisting for the most part of sulphuric acid thickened by
carbon.

11. From these phenomena Fourcroy and Vauquelin

Chap. IV.

A small quantity of ether is formed spontaneously, and without the assistance of hear, by the combination two parts of concentrated sulphuric acid and one part alcohol.

Theory of the decompourson of alcohol by sulphuric acid.

As soon as ether is formed, there is a production of ater at the same time; and while the first of these empositions takes place, the sulphuric acid undergoes change in its intimate nature.

As soon as the sulphurous acid appears, no more ther is formed, or at least very little; but the sweet all of wine passes over, together with water and acetic aid.

The sweet oil of wine having ceased to come over, othing further is obtained but the sulphurous and caronic acids, and at last sulphur, if the distillation be arried to dryness.

The operation of ether is therefore naturally divided to three periods: the first, in which a small quantity of ether and water are formed without the assistance of eat; the second, in which the whole of the ether which in be obtained is disengaged without the accompanient of sulphurous acid; and the third, in which the reet oil of wine, the acetous acid, the sulphurous acid, and the carbonic acid, are afforded. The three stages we no circumstance common to all but the continual fromation of water, which takes place during the whole the operation.

A combination of sulphuric acid and alcohol in equal ats does not boil at less than 207° of temperature, hile that of alcohol alone boils at 176°. Now since allition does not take place till the higher temperature,

it is clear that the alcohol is retained by the affinity of the sulphuric acid, which fixes it more considerably. Now organic bodies, or their immediate products, when exposed to a lively brisk hear, without the possibility of escaping speedily enough from its action, suffer a partial or total decomposition, according to the degree of temperature. Alcohol undergoes this last alteration when passed through an ignited tube of porcelair. The reason therefore why alcohol is not decomposed when it is submitted alone to heat in the ordinary apparatus for distillation is, that the temperature at which it rises in vapours is not capable of affecting the sepantion of its principles; but when it is fixed by the sulphuric acid or any other body, the elevated temperature it undergoes, without the possibility of disengagement from its combination, is sufficient to effect a commencement of decomposition, in which ether and water are formed, and carbon is deposited. Nothing more therefore happens to the alcohol in these circumstances than what takes place in the distillation of every other vegetable matter, in which water, oil, acid, and coal, are afforded.

Hence it may be conceived that the nature of the products of the decomposition of alcohol must vary according to the different degrees of heat; and this explains why at a certain period no more ether is formed but the sweet oil of wine and acetous acid. In fact, when the greatest quantity of alcohol has been changed into ether, the mixture becomes more dense, and the heat which it acquires previous to ebullition is more considerable. The affinity of the acid for alcohol being increased, the principles of this acid become se-

hydrogen, and forms much water, which is gradually vo- Chap. IV. latilized; while, on the other, the ether recaming a greater quantity of carbon, with which at that temperature it can rise, affords the sweet oil of wine. last ought therefore to be considered as an ether con--taining an extraordinary portion of carbon, which gives it more density, less volatility, and a lemon-yellow colou:.

Such is the ingenious explanation of the formation of sulphuric ether proposed by Fourcroy and Vauqueiin. They have succeeded completely in proving that the opinions formerly entertained respecting that singular process were erroneous; though their own explanation is not sufficiently precise to enable us to ascertain exactly the component parts of ether.

II. NITRIC ETHER.

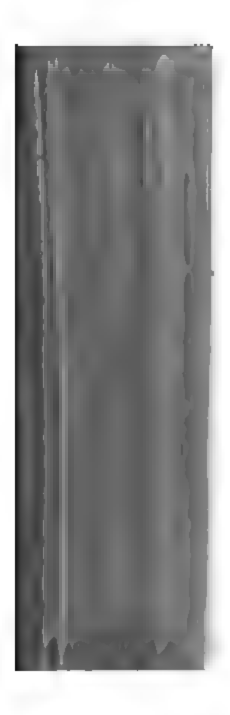
NITRIC ether is first mentioned in an epistle written by Kunkel to Voight, and published in 1681*; but no attention was paid to it by succeeding chemists till it was discovered a second time by Navier in 1742 +, and a third time by Sebastiani in 1746 t.

The method of preparing it, proposed by Navier, Preparawas this: Twelve parts of alcohol are put into a strong bottle, which is kept surrounded with water, or rather with ice: eight parts of nitric acid are poured in at intervals, the mixture being agitated after every addi-The bottle is then well corked, and the cork se-

Epistola contra Spiritum l'ini sine Acido.

¹ Diss. de Nitro, 1746. ± Mem. Par. 2742.

it is clear that the ...) LET. the sulphune . ". many because at the m Now organic for - धार त्रावा मध्य है expose! to a ign' . s. ct, the cosk is to be of escaping Ages " - quantity of nistings partial central ." or carry the ether un of temperature. or deawn; the whole when part ? rel, and by mean of a The reason of . . . sed to run out while sechen it as a conhezardous; for the qu params for " it rises in the segreat as often to but a in place of it a very tion of Allina - a glass phial the phurs ", " er this was poured & it ti fer e dechol was poured from 1 5 -2 of liquids in the ver ment of t L'echol uppermost, for Total ster. The acid and the water, and to than a dre without violence Special De ## · ! out risk. process for obtain. Bo but not attende of est mito a tubula ? e ed a luced, with part our hours, drop s the weight of and ether passe ide more militie == evening. The seed is to sof the Feto



gaseous form, while the liquid in to considered nitric ether, was a water, ether, nitrous, and acetic a led him to the following methor nitric ether.

Thenard's Process.

Equal weights of alcohol and m cific gravity 1-283, were put into beak of the retort was luted a glasged to the bottom of a long narrow with a saturated solution of com-From the top of this jar passed a went to the bottom of another simifilled with a solution of common a five similar jars were connected w half filled with a saturated soluti From the last a tube passed to a ceive the gaseous products in prothese five jars was surrounded with and salt to keep it as cool as post heat being applied to the retort a v It was found necessary to and even to moisten the outside of the in order to prevent the vessels fr

y. By this method, knowing the specific gravity of Chap. IV. vapour, and the products furnished by its decomposion, it would have been easy to have deduced its conmuents.

During the formation of nitric ether a vast quantity Gaseous gas is evolved. The Dutch chemists examined this and considered it as a compound of nitrous gas and ther; but Thenard has shown, that it is much more coplicated in its nature. The result of his examinafrom is, that it consists chiefly of nitrous oxide gas, mixed with a little nitrous gas, azote, carbonic acid, petic acid, nitrous acid, and a considerablet proportion

etherial vapour. But the proportion of this last in-

redient diminishes according to the degree of cold to

which it has been subjected.

products.

After the mixture of alcohol and nitric acid has Andresidue seased to give out ether, there remains in the retort bout three-fifths of the original quantity. This residue Thenard also subjected to examination. It has a yelnw colour and an acid taste. It consists chiefly of nater, holding in solution some nitric acid, some alcoa very small portion of acetic acid, and a matter which Thenard could not separate, but which very readily assumed the state of chargoal.

Thus it appears, that both the alcohol and acid are secomposed during the process of making nitric ether, and that the constituents of both enter into the compotion of the either formed. We are not sufficiently acmainted with the composition of alcohol, and with the exoportion of the other ingredients evolved, to be able a enter into the minutize of the decomposition. Some light may be thrown upon the subject by a careful exBock II. Division II. amination of the phenomena which take place carry the action of nitric acid on alcohol.

Actino of nitric sent on sicohol. 1. When equal parts of alcohol and nime soft is mixed, a violent effervescence takes place, spontage ously if the acid be concentrated; on the application the heat if the acid be diluted. This effers escence is one; to the emission of the gas, which the Dutch chemistic considered as a mixture of ether and nitrous gas, the which Thenard has shown to consist chiefly of entress oxide and ether. The Dutch chemists have called a nitrous etherized gas.

z. Nitrous etheriscid gra. This gashas a disagreeable ethereal odour: it bers with a yellow flame; is completely absorbed by want, alcohol, and the solution of potash; ammonia has to action on it. When fired along with oxygen gas it detonates. Sulphurie, sulphurous, nitric, and muratic acids, decompose it.

z. Ozalic acid. 2. When one part of alcohol and three parts of nitric acid, of the specific gravity 1.261, are mixed together, and a very moderate heat applied, a great quantity of gas is disengaged, which consists chiefly of nitrons either rised gas and nitrous gas. When only the part of the liquid remains in the retort, if it be allowed to cool, a number of crystals of exalic acid are formed to the process 1.107 parts of exalic acid may be obtained from 16 parts of alcohol 1.

Combustion of alcohol. 3. When one part of nitric seid is poured upon its own weight of alcohol, and one part of sulphure end is added a little after, the mixture takes fire and burns

Jeur. & Phys. zlv. 245.
 Sage, Jour. & Phys. 1, 346.

† Scheele and Hermhstadt.

ith great rapidity. When this experiment is perormed in close vessels, the products are ether and oil, lesides what remains in the vessel in which the com-Instion takes place ".

4. When nitric acid, partly saturated with mercury, Howard's Is poured upon alcohol, and heat applied, the products fulmined we nearly the same, but the phenomena are very diffeent. The curious appearances which accompany this mixture were first observed and explained by Mr Howardt. The process, as described by him, is as follows: Dissolve, by means of heat, 100 grains of percury in a measured ounce and a half of nitric acid, If the specific gravity of about 1.3. Pour this solution pon two measured ounces of alcohol, and apply heat the mixture begins to effervesce. The heat is then be withdrawn. The action becomes violent, and conloues for some time; a dense white smoke issues from he vessel, which is heavier than the atmospheric air. ind may be poured into glass jars, where it continues for some time like fine white clouds. Mr Howard has made it probable that this fume is composed of etheused nitrous gas holding oxide of mercury in solution. Meanwhile a white powder falls to the bottom of the pixture. When the effervescence is over, this white pwder is to be separated by filtration, washed with rate water, and dried in a heat not exceeding 212°.

Mr Howard has examined the properties of this powder, which has the appearance of minute crystals. He has given it the name of fulminating mercury.

P Brugnatelli, Ann. de Chim. xxis. 327.

[!] Nicholson's Journal, iv. 173.

of mercury, and more of the peculiar vegetable matter. Chap. IV. When the mixture is boiled for half an hour, the powder is composed of oxalate of mercury and a very small quantity of vegetable matter. It does not detonate, but = decrepitates when heated *. These experiments of Fourcroy enable us to reconcile the seemingly opposite results of Howard and Berthollet +.

III. MURIATIC ETHER.

AFTER the discovery of sulphuric and nitric ethers, various attempts were made to obtain ether by the action of muriatic acid on alcohol: But this acid in its usual state is too much diluted with water to act with much energy upon alcohol. It was thought necessary, therefore, in order to procure muriatic ether, to employ the acid in a different state. Two methods have been discovered.

1. Those muriatic salts are chosen which may be obtained dry, and at the same time have a strong affinity for water. All the salts which have been hitherto tried 1. By sales with success have a metallic base in the state of a peroxide 1. The oxymuriates of mercury, iron, arsenic, and antimony, produce ether when distilled with alcohol: but the salt which answers best is the oxymuriate

Prepara

[●] Journal of the Royal Instit. i. 256.

[†] According to Berthollet, fulminating mercury is composed of ammonia, oxide of mercury, and altered alcohol, which produces carbonic acid when decomposed. Pbil. Mag. xii. 92.

I I distinguish this state of oxidizement in the metallic salts by prefixing exy to the usual names of the salt.

Book II. Division II. of tin. By means of this salt Continues in ether, in 1750, by the following process Turns of furning oxymuriate of tin and one part of action mixed together; and after the vapours and her salt ced have subsided, the mixture is put into a react which two large receivers are attached, and distributed two large receivers are attached, and distributed the comes over first a little alcohol, then thereof. This salt has been lately recommended as very pay for making muriatic other by Klaproth t.

a. By the

g. Pure alcohol is sa urated with municipal free from water as possible. The following what mula recommended by Mr Basse. Keep a quantition salt for an hour in a state of fusion. It to deprive it of its water of crystallization, parts of this salt into a tubulated retort, to the which is fitted a bent tube, plunging into Woulf the, containing 10 parts of alcohol as strong as parts.

Introduce into the retort, in small quantities at 10 parts of the most concentrated sulphuric at lowing the common air to escape from the bott taining the alcohol, then distil in a sand bath murtaire acid comes over, keeping the alcohol cool as possible during the process. The alcohol saturated with acid, is put into a retort, and one it distilled over. Agitate this portion with an ales, and then decant off the ether which swims burtace it usually amounts to 2½ parts I.

This process was discovered for the process was discovered for the first p.

¹ Lite + Amount 1 195 1 99.

^{. -} The paper of party property water and it to the

by little was known of the properties of muriatic till Gehlen published a dissertation on the sub-1804 . He employed two processes: 1. the of the furning oxymuriate of tin on alcohol; 2. occas of Basse detailed above. Both of them furmuriatic ether, the peculiar properties of which a described with accuracy. Thenard published Lissertations on it in 1807 +, pointed out the simscocess for obtaining it, examined the effect of oxytes on alcohol, described the properties of muciher in detail, and made a set of experiments to ain its constituents. To the dissertations of these emists we are indebted for all that we know of ry remarkable substance.

process recommended by Thenard for procuring Thenard's her is the following: Equal bulks of muriatic acid phol, both as strong as possible, are put into a retort. a size as not much more than to hold the mixture. grains of sand should be put into the retort, to preeviolent boiling which might otherwise take place. the beak of the retort a tube passes into a glass ice the size of the retort, and furnished with three . This jar should be half filled with water, of

reperature of about 70°. Into the second mouth

smic acid, by distilling a mixture of two parts of common salt are of sulphure acid into a Woulfe's apparatus containing two Leohol. Min the sat rated alcohol with half a part of black manganese, and put into the Woulfe's apparatus a solution of when water, and destil with a low heat. The ether and oxyacid pass over, and the potash prevents the acid from acting so the other. See Ann. de Chim. EXE'V. 141.

† Mem. d' Arauel, i. 115 149. 357. Men's Jour. 11, 206.

Book II. Division II.

of tin. By means of this sale ether, in 1759, by the follow of fuming oxymuriate of tin mixed together; and after ced have subsided, the newhich two large receive. There comes over first. This salt has been lanfor making muriatie.

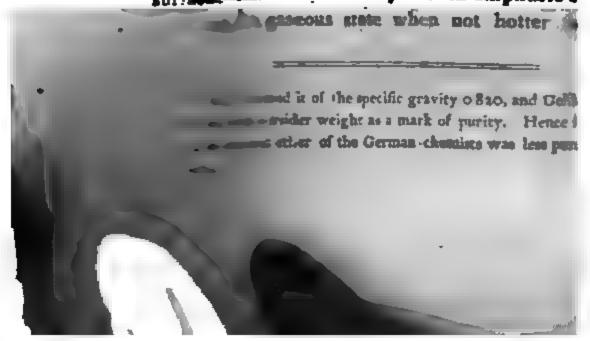
2. By the acid.

2. Pure alcohol
free from water a
mula recommen
common salt fo
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air being 1.
ives its own hell

. 4 66

Introduction cold of 52°, it loses it lowing the cold of 52°, it loses it lowing the cold of 52°, it loses it taining the cold as the state by passing it into a dry muriation. On the water, very liquid, has no a saturation cold as the same smell and the it distributed state. At the temperature of 41°, ley, many of the specific gravity 0.874°. It surfaces, satisfaces than alcohol, or even sulphuric to



the presence of any Chap. IV. vetable blues, nor line ley, or oc-'d with niwith a ured acid This Gehlen: .rd. When ontact with an uriatic acid, and atrate of silver after the quantity of preci-But in neither case is prived of the property of sien burnt.

of muriatic ether no gaseous prolived but muriatic ether; nor is any
mee evolved, unless a portion of water
d. A portion of the muriatic acid as
me alcohol disappears; and when the ether
apposed, exactly the portion of acid is evolved
which had disappeared. It is extremely difficult from
the facts to form an accurate notion of the way in
which the ether is formed. Is it a compound of alcohol and muriatic acid?—The little effect which these
two bodies have on each other, even when mixed in
the state of vapour, renders that opinion unlikely. It
is equally difficult to conceive the state of the muriatic

acid in that liquid. It seems to be perfectly neutrali-

med, as all its usual properties are concealed; and it must

is of Thenard it appears, that Composi-

Book H. Diwana H. stances on which it acts with the greatest energy indicate its presence when mixed with the other. At the same time, difficult as it is to explain how it is controlled, it is more probable that it exists in the state of marriatic acid than decomposed, if we consider how difficult it is to decompose this acid, and how obstinately a bas resisted the numerous attempts to ascertain its composition.

> 36.61 carbon 23.31 oxygen 10.64 hydrogen

If any confidence can be put in this analysis and in that of alcohol by Saussure, we may infer from them, that muriatic ether does not contain the alcohol in the state of alcohol, since the carbon, oxygen, and hydrogen to not bear to each other the same proportion in the e-

Mem. D'Aravil, 1. 341.

ther that they do in alcohol. The proportion of car- Chap. IV. bon in the ether is a good deal more, and that of the oxygen less than in the alcohol.

After the discovery of oxymuriatic acid, Scheele Action of showed that ether might be obtained by distilling a new ton mixture of alcohol, black oxide of manganese, and muriatic acid; or by distilling sulphuric acid, common salt, black oxide of manganese, and alcohol: but the quantity which can be obtained by this process is trifling. for the oxymuriatic acid acts upon the ether formed, and converts it into a kind of oil. Indeed, if we believe Mr Basse, ether is never obtained by means of oxymuriatic acid, but merely an oil which sinks in water. This has been amply confirmed by the late experiments of Thenard +; from which we may conclude, that exymuriatic acid converts alcohol into oil, and not into ether, and that which has been taken for ether is nothing else than alcohol holding some of this oil in solution.

orymuris-

IV. ACETIC ETHER.

ETHER may be produced also by the action of acetic acid on alcohol. This was discovered by the Count de Lauraguais in 1759 1. He obtained it by distilling a mixture of acetic acid and alcohol with the same precautions as are employed in the distillation of sulphuric ether.

The process, as corrected by Pelletier, is as follows: Prepara-Mix together, in a retort, equal quantities of acetic tion. acid (from acetate of copper) and alcohol, and distil over the alcohol. Pour it back into the retort, and di-

[&]quot; Jour de Chim, iv. 88. + Mem. D'Aremil. t. 47. 1 Just de Scau. 1759, p 324.

a violent boiling, the phosphoric ek, and striæ in abundance appeared on ak of the retort. The distillation was phosphoric acid became dry. There the receiver, 1st, 120 parts of alcohol, of ether; 2d, 260 parts of a colourless light etrong of ether; 3d, 60 parts of watersaer, over which swam 4 parts of a yellow th resembling the sweet oil of wine in apanother liquid of a disagreeable odour, egetable blues. When saturated with porated, it left a quantity of acetate of a. The lime water became milky, but end of the process. A quantity of gas bich burnt like ether, and seemed to conher uncondensed.

products being rectified on muriate of Proportion bo parts of a liquor bearing the closest sulphuric ether. It had the same smell me specific gravity, dissolved in 8 or 10 ater, boiled at the temperature of 100°, and phosphorus, burnt with a white trace of charcoal, but giving no indicaonce of any acid.

ears that phosphoric ether approaches alphuric ether, if it be not absolutely s, and differs very considerably from niad acetic ethers.

be formed also by several other acids. d it by distilling a mixture of fluor spar, anganese, alcohol, and sulphuric acid; means of oxalic acid. Scheele found

Division II.

that the following acids did not form ein hol .

- 1. Mariatic,
- 4. Benzoic,
- 2. Fluoric,
- 5. Tartaric,
- 3. Boracie.
- 6. Citric,

Thus it appears that there are various specific thers differing very much from each other is perties. Sulphuric ether is the lightest but volatile of the whole. In muriatic and active acid seems to enter as a constituent part, but its page are concealed altogether while the liquid remain composed. It is not unlikely that nitric ether a so contain an acid. It is obvious that the theory formation of nitric, muriatic, and acetic ethen: quite different from that of sulphuric and phosph thers. The theory of the two last is probablys

SECT. III.

OF VOLATILE OILS.

THE term oil is applied to a number of unct quids, which, when dropt upon paper, sink inte make it seem semitransparent, or give it what i a greasy stain. These bodies are very numero have been in common use from time immemoria mists have divided them into two classes; name latile and fixed oils. We shall consider the prope

Scheele, ii. 117.

If these classes in this Section. The fixed oils by our attention in the next. A third class wht be added which possess intermediate proween the fixed and the volatile.

Chap. IV.

THE OILS, called also essential oils, are distiny the following properties:

uid; often almost as liquid as water; some- Characters.

y combustible.

scrid taste and a strong fragrant odour. tatilized at a temperature not higher than

sble in alcohol, and imperfectly in water. aporate without leaving any stain on paper. Is last test it is easy to discover whether they adulterated with any of the fixed oils. Let of the volattle oil fall upon a sheet of writing d then apply a gentle heat to it. If it evapothout leaving any stain upon the paper, the oil but if it leaves a stain, it has been contaminaname fixed oil or other.

le oils are almost all obtained from vegetables, Preparaexist in every part of plants; the root, the tion. wood, the leaves, the flower, and even the igh they are never found in the substance of dons; whereas the fixed oils, on the contrary, always contained in these bodies ".

the volattle oils are contained in great abun-Mants, they are sometimes obtained by simple . This is the case with the oil of oranges,

Fourcray, vii. 352

bar Impo

Caches see it bety amount but in genture in a community in our application. The first cap of community in our application of by the applications in accordance of by the applications in accordance of by the applications in accordance of an outcomes over along with the accordance of the outcomes over along with the product and application that out or perpetition in the because and community of the outcomes. Under a strong of the Caches and a personal of residues bothers. This is the accordance with our or the pentition. White is one of Caches, a reside to residues party, caused improve the caused tripers.

Propertie

totalin our accessorangly numerous. They are some our now enount, but as their use in chemisty self-innece they have not intheric need. Subjected as mechanic mechanic myestigation. They differ grant their proper as from each other. but it is impossible as present to, we a detailed account of each

League e e

many more are a impic at water, who have not of time appearance which we usualty considers a trapearance which the following mamery, of time of the only viscidity. It varies in them, in the property of mass, considers, common, sustaines, cloves, combanon. Others have made, assumes, cloves, combanon. Others have property of becoming solid. This is the case with the others have made of paralley, fermel, amseed, balan. Others my made by slow evaporation. This is the case with the time of the case with the others my made of paralley, fermel, amseed, balan. Others my made by slow evaporation. This is the case with the thy slow evaporation. This is the case with the thy slow evaporation. This is the case with the thyme, peppermint, marjoram. The oil of name

maily the consistence of butter . This is the aso with the oil of hops and of pepper.

The colour of the volatile oils is as various as Colour-

other properties. A great number are limpid Mourless, as oils of turpentine, lavender, rosemary, 💃 aniseed. Some are yellow, as spike, berga-Some are brown, as thyme, savony, worm-Others blue, as camomile, motherwort. Others as milfoil, pepper, hops, parsley, wormwood, it, juniper, sage, valerian. Others, though at colourless, become yellow or brown by age, as , cinnamon, sassafras †.

Their odours are so various as to defy all des- Odour. in. It is sufficient to say, that all the fragrance vegetable kingdom resides in the volatile oils. taste is always acrid, hot, and exceedingly un-

Their specific gravity varies very considerably, Specificant aly in different oils, but even in the same oil in dif- vity. circumstances. The following are the specific des of several of the volatile oils, as ascertained by

fsassafras1.094	Oil of Mint 1975
Cinnamon 1:035	Numegs '948
Cloves 1 '034	tansy 946
Fennel 1997	Carraway seeds 940
Dill 904	Origanum940
Penny royal 978	Spike936
Cummin. 1975	Rosemary 934

roy, vii. 505. Eng. Trans.

† Neuman's Chem. p. 272.

Street E.

Oil of Juniper berries '611 Oil of Turpes Oranges.....'888

Printer.

when the volatile oils are heated in the they evaporate readily, and without alterant their peculiar odours all around; but there detable difference between the different out pect. When distilled in close vessels, they readily assume the form of vapour. Hence their odour, become darker in colour, and a decomposed. Oils do not seem very suscept suming the gaseous form, unless some other as water, be present.

When exposed to the action of cold, they estate fixed oils; but the temperature necessary whis effect varies according to the oil. Some as oil of anise and of femal, become solid at perature of 50° I frezen oil of bergamorte and become liquid at 23° and of turpentine at 14° queron exposed several volatile oils to a cold. They congealed or rather crystallized partially the same time emitted an elastic fluid. These consisted partly of the oils themselves, partly substances. Some of them had the properties zoic acid †.

Charges by

o. Volatile oils, when exposed to the action close vessels, and excluded from common six very singular changes. Their colour become they acquire a great deal of consistency, and their gravity is considerably increased. The

Margueron, Jour. de Phys. 2lv. 136.

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shanges is but imperfectly known. Tingry, to we are indebted for these interesting researches. goved that light is a necessary agent. It was supformerly that they were occasioned by the abof oxygen; and when oxygen is present, it en ascertained that it is absorbed: but Tingry oved that the same changes go on when oxygen aded. This philosopher ascribes them to the of light. If this be the real cause, the quantity he fixed must be enormous; for as the specific of the oils is increased considerably while the continues the same, it is evident that the absolute must be increased proportionably. One cirince, however, renders this conclusion somewhat at least in its full extent; and that is, that the y of change was always proportional to the of the oil and the quantity of air contained in hael .

Absorb oxygen,

When volatile oils are exposed to the open air, they lly become deeper coloured, and acquire more re viscidity, while at the same time their odour hes. Dr Priestley first ascertained that they oxygen with rapidity, and that the changes are this absorption. He tried the experiment only of turpentine, but he found that the air above aint and cinnamon, confined in phials half full, rived of its oxygen †. He ascertained likewise, dependent of this disposition to absorb oxygen, arpentine has the property of imbibing a consi-

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[.] Tingry, Jour. de Plys. alvi 16., and 249.

¹ Pricatley on de, 11 232.



The appeare the volatile oils assume.

Then the volatile oils are heated
the they take fire and burn w
thing a vast quantity of smo
timbustion, besides the soot, a

When agitated with water, the provide oils render it n ilky, and we emiliar odour. Several of then in fithe sugar be afterwards disconnected as a manner solution, to which are m has been given. Marguer to of the oils.

The action of the simple combi-

brogen as far as is known, neither are they altered Chap. IV. charcoal. When digested upon sulphur at the temtrature at which the sulphur melts, they dissolve a attion of it, acquire a brown colour, and a disagreetaste and smell. These preparations are call-Calsams of sulphur. A portion of the sulphur crys-Mizes as they cool . When these balsams are heated Fongly, a vast quantity of gas (probably sulphureted drogen) is evolved so rapidly as to occasion very plent explosions, unless proper precautions be taken †. The volatile oils dissolve likewise a portion of phosfocus in a digesting heat; but most of them deposite whole of it again as the solution cools. pinted out a method of rendering the solution permaant. It was the following: Triturate together ten larts of camphor and one of phosphorus. This mix-Fore dissolves readily in most volatile oils, as in oil of hoves, and forms a solution which has the property of endering every thing luminous which is subbed with It, and this without combustion t. This seems to have seen the solution so much used by Boyle, under the some of liquid phosphorus.

10. The alkalies and earths act but feebly upon the Of alkalies polatile oils. The French chemists have proposed to and earths, ive the combinations which these bodies form with he volatile oils the name of savonules, which Dr Pear-

Element de Chymie of the Dijon Academy, ich 357.

I Hoffman relates a remarkable story of the violent effects of such an apionus by way of caution to the chemists of his time, -Observationes Mys. Ches. p. 308.

[|] Halman, Obiero. Phys. Chem. 5. 307.

tate treated as above to the and a safe wax, soluble in the trular a tecomposed by an alandary of the units readily with the at a tecomposed by an alandary of the analysis at it has approached the analysis.

... acid is thrown upon them suddenly, at ... rand state, it acts with such energy as to so ... but when sufficiently diluted with water ... tem, and converts them into a yellow solution Oxymuriatic acid acts in the ... though with less energy.

Let ned with care, but it cannot be remarkable.

Let ned with care, but it cannot be remarkable.

Let has tried the effect of some of the sales of the sale is gradulated over nitrate of mercury, the sale is gradulated of mercury, in like manner, deepens the sale increases the consistence of oils of cinon, the sale is at the same time partly converted into murlate of mercury. Neither the muriate of mercury, not

[.] Joer. & Pigs. avi. 409

[†] Elemens de Chymie of the Dijun Academy in juice

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the sulphuret of that metal, produce any change in the oils of lavender and rosemary; but by this last oil the red oxide of mercury is converted into the black, though the oil does not experience any sensible change. The oxymuriate of antimony is likewise decomposed by the oil of rosemary *.

- 13. From the effects of the acid supporters on the volatile oils, and from the products which they yield when burnt, it has been concluded that they are composed of hydrogen and carbon, sometimes united with various proportions of oxygen according to circumstances: but no exact analysis has yet been made of any of them.
- 14. Volatile oils are applied to a great number of uses: Some of them are employed in medicine; some of them, as oil of turpentine, are much used to dissolve resins, which are afterwards employed as varnishes. Not to mention their employment in painting and in perfumery.

SECT. IV.

OF FIXED OILS.

The fixed oils, which are of such extensive utility in Discovery. the arts, were known at a very remote period. They are mentioned in Genesis, and during the time of Abraham were even used in lamps †. The olive was very early cultivated, and oil extracted from it, in Egypt.

Ann. de Chim. xlvii. 66.

† Gen. 14. 17.

ally deposi-Chap. IV. so in the eggs

in several partiarticulars in comall the fixed oils is heir differences to accompletely ascertained, .erto been made; but it coils hitherto tried have e products. In the present t would be useless to give a all the fixed oils, as even the .m have not been accurately as-

ually a liquid with a certain degree Properties. ing to the sides of the glass vessels in ained, and forming streaks. ransparent, having always a certain de-, most usually it is yellowish or green-: is sweet, or nearly insipid. When fresh or no smell.

.st also in the vegetable kingdom a considerr of bodies, which at the ordinary temperatmosphere are solid, and have hitherto been Palm oil may be mentioned s fixed oils. ple, which has been lately subjected to a amination by Dr Bostock †. The various sed in India and Africa as substitutes for as unquents, may likewise be mentioned.

[₩]y, vii. 319. MAN Journal, Xvi. 161.

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Most of them are obtained from the seeds of trees; different species of the bassia, as the butyracea, longifolia, latifolia, obovata, yield this butyraceous matter. They have been described by Dr Roxburgh . The shea or butter tree of Africa described by Park, seems also to be a species of bassia. These substances, from the experiments of Dr Bostock, appear to differ a little from the liquid fixed oils in their properties, and to approach the nature of wax. Thus they are sensibly soluble in alcohol and ether, and do not combine so readily with alkalies as the fixed oils.

Specificgra-

2. All the fixed oils hitherto examined are lighter than water; but they differ greatly from one another in specific gravity. The same difference is observable in different samples of the same oil. The following Table contains the specific gravity of such oils as have been examined.

Oil of palm*...968

Hazel-nuts*941

Poppies*..939

Linseed†..932

Almonds* 932

Walnuts*..923 to 947

Beech-nut* 923

Ben*....917

Olives†...913

Rape-seed† 913

Cacao‡...892

Action of hose.

3. Fixed oil does not begin to evaporate till it be

[§] Nicholson's Jour. xix. 372.

^{*} Fabroni, Crell's Annals, 1797, iL 123.

[†] Shaw's Boyle, ii. 346.

[‡] Brisson

heated above the boiling point of water. As the heat increases, a pretty copious vapour may be seen rising from it; but the oil does not begin to boil till it is heated nearly up to the temperature of 6000. At that temperature it may be distilled over; but it is always somewhat altered by the process. Some water and acetic acid seem to be formed, a little charcoal remains in the retort, and the oil obtained is lighter, more fluid, and has a stronger taste than before. Oil, thus distilled, was formerly distinguished by the name of philosophisal oil. During the distillation, a great quantity of heavy inflammable air is obtained.

Fixed oil, when in the state of vapour, takes fire on the approach of an ignited body, and burns with a yellowish white flame. It is upon this principle that candles and lamps burn. The tallow or oil is first conwerted into the state of vapour in the wick; it then takes fire, and supplies a sufficient quantity of heat to convert more oil into vapour; and this process goes on while any oil remains. The wick is necessary to present a sufficiently small quantity of oil at once for the heat to act upon. If the heat were sufficiently great to keep the whole oil at the temperature of 600°, no wick would be necessary, as is obvious from oil catching fire spontaneously when it has been raised to that temperature. When oil is burnt in this manner, either in the open air or in contact with oxygen gas, the only new products obtained are water and carbonic acid.

When exposed to the action of cold, fixed oils lose their fluidity, and are converted into ice; but this change varies exceedingly in different oils.

3. When fixed oils are exposed to the open air or to Action of oxygen gas, they undergo different changes according to

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Most of them are ferent species of latifolia, obovat. have been desc butter tree of the a species of periments of the liquid fitthe nature of alcohol and alkalies as

of absorbing oxygen;

of absorbing oxygen;

me more and more vissolid state, being appaNow there are some
mency after they have bemency after they have bemency and assume
rwax. This circumstance
rwax. This circumstance
rwax is in fixed oils into

. :hat become opaque are called

Spicalegra-

2. All than water specific gradifferent contains cxamino

re used as the vehicle of paints and and hemp seed oils, These oils in their natural state grof drying oils but imperfectly. To . whe use of the painter and varnish armioiled for some time in an iron pot. me men and of carbureted hydrogen gas is They become deeper coloured, rer consistency. It is common for some es is them on five, to allow them to burn for = rexinguish them by covering up the vesme they are contained, and to continue the in acquire the proper degree of viscidity. * mass they lose their unctuous quality in a saste, so as not to leave a greasy stain upon approach the nature of resine, with this difin they do not become brittle, but retain a was a soughness and ductivity, not unlike what in is called wrought rosin, or shoemaker's rohis common also in preparing the drying oils to

. Iction of heat.

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The litharge. The change which y this process has not been pre-Probably they absorb oxygen from ow that they undergo a partial denen they burn for some time, their is much more completely destroyed ethod which has yet been practised. lowed frequently in preparing the dryvarnishes, and always for printers ink, es to be as free as possible from all unc-

sas been found preferable to all other oils for k; though the dark colour which it acquires iling renders it not so proper for red ink as Linseed oil is considered as next after nuthis respect. Other oils cannot be employed, they cannot be sufficiently freed from their osity. Ink made with them would be apt to e off and smear the paper while in the hands of the k-binder, or even to spread beyond the mark of the pes, and stain the paper yellow. The process for aking printers ink is as follows:

The oil is made to boil in an iron pot only half Printers filled, set on fire, and allowed to burn for half an hour or more, then boiled gently till it acquires the proper consistence. In this state it is called the varnish. kinds are prepared, a thicker and a thinner. thicker is of such a consistence that it draws into threads when cold like weak glue. This varnish is afterwards ground with lamp-black in the proportion of two ounces and a half to sixteen ounces of oil. When newly prepared oil is used for making ink, it is said to be necessary to add a little boiled oil of turpentine and a little

Book II. Divis en II. litharge; but this is said to have the effect of causing the ink to stick so firmly to the types, that it can wat difficulty be removed. Old oil does not require the addition.

Oil prepared by the process above described, is said insoluble in alcohol and water, but it unites readily to more oil. It dries into a tough mass like turpentury, and afterwards is scarcely susceptible of uniting with of Dr Lewis found that linseed oil, when thus converted into a thick varnish, lost 4th of its weight; when boiled till it became quite stiff when cold, it lost nearly; of its weight. The property which printers ink his of adhering to moistened paper shows that the oily return of the body is greatly altered. In some tespects at has approached the nature of mucilage, though in other the difference is very great.

Fat oils.

5. The fat oils, when exposed to the atmosphere, gradually become thick, opaque, and white, and assume an appearance very much resembling wax or tallow. Olive oil, oil of sweet almonds, of rape-seed, and of bea, belong to this class.

When oil is poured upon water, so as to form a that layer on its surface, and is in that manner exposed to the atmosphere, these changes are produced much sometre. Berthollet, who first examined these phenomena with attention, ascribed them to the action of light but Sennebier observed that no such change was produced on the oil though ever so long exposed to the light, provided atmospherical air was excluded; but that it took place on the admission of oxygen gas, whether the

[&]quot; Lewis, Phil. Cept p 327.

was exposed to the light or not ". It cannot be Chap. IV. coubted, then, that it is owing to the action of oxygen. is supposed at present to be the consequence of the timple absorption of oxygen and its combination with the fat oils.

6. The action of fixed oils upon the simple combus- Action of ables is not very remarkable.

simple combustibles.

Hydrogen, as far as known, does not act upon them. When they are filtered through charcoal powder, they are rendered purer; but on account of the great difficulby of separating the charcoal from fixed oils, it cannot be employed with advantage for purifying them †. Black paint is usually nothing else than charcoal in some state or other ground up with a drying oil.

The fixed oils likewise dissolve a small proportion of phosphorus when assisted by heat. The combination succeeds easiest if a mixture of oil, water, and phosphorus be boiled for a little in a glass vessel. These oily phosphurets emit the odour of phosphureted hydrogen, and yield, when distilled, a portion of that gas. When rubbed in the open air, or when spread upon the parface of other bodies, they appear luminous, in consequence of the combustion of the phosphorus. When hot oils saturated with phosphorus are allowed to cool, the phosphorus crystallizes in octahedrons, as Pelletier scertained.

They readily dissolve sulphur when assisted by heat. The solution assumes a reddish colour. When distilled, there comes over a great quantity of sulphureted hy-

[·] Ann de China. 21, 89.

[†] Kels, Crell's Annals, iin 274. Engl. Trans.

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solution is allowed to col, to exystals. By this process this are regular octahedrons.

tre all insoluble in water. When the mixture becomes may the mixture becomes may the correspondent and swim upon the correspondent of a mucilaginous substant,

eresents the oil from separating, and one are a permanent milkiness. Such make the ends of the formed to ends, as almonds, with water, the two eresent to form an emulsion, oil and make the eresent to the seed.

the in it even after being thickened by boldnes be united to an alkali, and afterwirds an acid, they will be found to have acrespectly of dissolving in alcohol! a preof has undergone an alteration in its composquented to an alkali.

with each other, with volatile oils, and

mportant compounds called soaps. The fat we these combinations more readily than the The earths likewise combine with these and form a kind of soap insoluble in water-fore not capable of being applied to the common soap.

. Same de Layane of the Dijon Academy, sii. 400.

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.5 is known at preand ther has the muriatic Chosphoric acid, when tell air, and gives them a peassisted by heat: a proof The sulphuric acid acts with the oils become immediately mily the properties of bitumen minuance of the action. If the main long enough, they are com-. water is formed, charcoal precipi-: evolved †. Other products doubtless . appearance, though the action of this .as not yet been examined with sufficient acid acts with still greater energy. When andenly upon the drying oils, it sets them on The same effect is produced upon the fat oils, ed the acid be mixed previously with a portion sulphuric. When the nitric acid is sufficiently l, it converts the drying oils into a yellow resike mass, and the fat oils to a substance very like But the action of this acid upon fixed oils has

t been sufficiently examined.

ny attempts have been made by chemists to form Acid soaps. nent compounds of the concentrated acids and oils the name of acid soaps. The only acid which and to answer was the sulphuric. Achard puba number of experiments on these compounds. dissolve in water, and lather like common soap;

Elemens de Chemie of the Dijon Academy, iii. 142-

[†] Fourcroy, vii. 330.

ence of an ointwater decanted off. the consistence of a Sweet pri Scheele termed the t crystallize, is soluconverted into oxalic When heated, it is n oil, and partly volatiace he obtained also from from oil of almonds. Even pyielded him a little of it *. pposed to exist in all fixed eir rancidity by putrefaction. ne degree diminished by agitacompletely destroyed. Mr Dositation with the fixed alkaline sole answers the purpose sufficiently ls for burning in lamps; but that erty of coagulating a portion of the ver may be prevented by adding a

oil is burnt, the only products arec arbonic Composi-.er. When repeatedly distilled, or when ugh a red hot tube, it appears to be comwerted into water, carbonic acid, and heavy ble air. Lavoisier analysed olive oil by burna vessel filled with oxygen gas. During the on there was consumed

odies from the oil +.

g brine, which occasions the separation

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ciple of oi

tion of oils.

le's Opens. ii. 189.

† Nicholson's Jour. v. 5.

Book II. Division II. Total......66.65

The products were carbonic acid and water. The case bonic acid obtained amounted to 44'50 grains in weight of the water could not be accurately ascernated; but as the whole of the substances consumed use converted into carbonic acid gas and water, it is evident, that if the weight of the carbonic acid be too tracted from the weight of these substances, there mist remain precisely the weight of the water. Mr Lavoseier accordingly concluded by calculation, that he weight of the water was 22'15 grains. Now the quality of oxygen in 44'50 grains of carbonic acid gat a 32'04 grains, and the oxygen in 22'15 grains of wite is 18'82 grains; both of which taken together amount to 50'86 grains, precisely the weight of the oxygen gas employed.

The quantity of charcoal in 44.50 grains of carbonic acid gas is 12.47 grains; and the quantity of hydrogra in 22.15 grains of water is 3.32 grains; both of when, when taken together, amount to 15.79 grains, which is the weight of the oil consumed.

It follows, therefore, from this analysis, that 15:78 grains of oil are composed of 12:47 carbon

3:32 hydrogen

Olive oil therefore is composed of about

79 carbon

21 hydrogen

100

[.] Men. Por. 1724, and Jour. de Phys. for 1787. July.

This, however, can only be considered as a very im- Chap. IV. perfect approximation towards the truth. The fixed oils no doubt vary in the proportion of their constituents; and the phenomena of their decomposition lead us to conclude, that many of them contain oxygen as a component part .

BESIDES the volatile and fixed oils, there is another Poisonous set which exist pretty frequently in the vegetable king-Their chemical properties are intermediate between the fixed and the volatile oils. Like the volatile oils they dissolve in alcohol; but like the fixed they cannot be distilled over with that liquid. Hence they may be obtained by digesting the vegetable substance that contains them in alcohol, and then separating the alcohol from the oil by distillation. They have all a strong acrid taste, and most of them possess poisonous qualities. The alcoholic solution of some of them has the property of precipitating sulphate of iron of a reddish colour, which becomes green when mixed with an alkaline solution. Vauquelin detected an oil of this nature in the root of the helleborus hyemalis †. A similar oil appears to exist in tobacco and in many other plants.

Besides the oils which exist ready formed in the vegetable and animal kingdom, there are a variety of others which are obtained when animal or vegetable bodies are distilled by ments of a heat above that of boiling water. These oils have received the appellat of of repyrounder, because they are formed by the action of fire. They have n ver bein examined with attention; but most of them seem to possess the propersies of the volatile oils. Their odour is always exceedingly disagreeable, and their taste acrid.

⁺ Ann. de Mus. & Hist. Nat. No. 2lui. 89.

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SECT. IV.

OF BITUMENS.

The term bitumen has often been applied by chemisa to all the inflammable substances that occur in the earth; but this use of the word is now so far limited, that religious and mellite are most commonly excluded. It would be proper to exclude amber likewise, and to apply the term to those fossil bodies only which have a certain resemblance to oily and resinous substances. In this restricted sense the word is used in the present Section.

Bituminous substances may be subdivided into two classes; namely, bituminous oils, and bitumens, properly so called. The first set possess nearly the properties of volatile oils, and ought in strict propriety to be classed with these bodies; but as the chemical properties of bitumens have not yet been investigated with much precision, it was deemed rather premature to separate them from each other. The second set possess properties peculiar to themselves. Let us endeavour to describe the substances belonging to these two classes at far as possible.

I. BITUMINOUS OILS.

ONLY two species of bituminous oils have been his

[Name

mentioned, but their existence has not been sufficiently Chip. IV. murben soused. These two species are called p eroleum, must multha or sea-wax; the first is liquid, the second moliu.

1. Petroleum is an oil of a brownish yellow colcur. Petroleum. When pure, it is fluid as water, and v ry volatile ". Its specific gravity valies from 0.7-0 to 0.878 t. It has a pecu ist smell. When heated, it may be distilled over without alteration. It will s with alcohol, ether, voiatile and fixed only, and, as her as known, possesses all the characters of volatile oils

Petroleum is found in the earth in various states of parity; sometimes without any mixture of foreign substances. In this state it is usually distinguished by the name of naphtha, and is said to occur in great abundance on the shores of the Caspian and in Persia. It occurs also in different parts of Europe, especially Italy and Germany. When less fluid and darker coloured, it is commonly called petroleum. It is supposed to owe this increased spissitude and deepened colour to the action of the air. When distilled, it yields a quantity of pure petroleum, while a portion of bitumen of the consistence of tar or pitch remains behind. When long exposed to the air, petroleum becomes black, and acquires the semifluidity of tar. In this state the greatest part of it is insoluble in alcohol; so that it has assumed the state of true bitumen.

The volatility seems to have been over-rated by older chemical writers. On susing well rectified petroleum with water in a retort, I found that the water could be read by distilled over in a moderate heat, while the whole petroleum remained behind.

[†] See Kirwan's Minerangy, it. 42.

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and for lamps. It is employed also as a minous bodies, and of the proper bimmens, and the

ne Baikai lake in Siberia. It is white, meliheated, and on cooling assumes the consistence of
herace. It dissolves readily in alcohol, and is
espects appears to possess the characters of a silande oil. Its properties, however, have been a
perfectly examined. Kimproth found that
substance was obtained by distilling a species
coal, called earth coal by the Germans.

Steeners-

The substance described by Kirwan under the substance described by Kirwan under the substance tallow, and said to have been found on of Finland, in the Swedish lakes, and in a four Strasburgh, seems to approach very nearly to Its specific gravity is 0.770. It is white, britt paper like oil, melts when heated, and burn blue flume and much smoke; dissolves in hot alcohol, but readily in olive oil †.

II. PROPER BITUMENS.

THE true bituminous substances may be dis

- 1. They are either solid or of the consistence
- 2. Their colour is usually brown or black.
- 3. They have a peculiar smell, or at least

Bestrage, ili. 305.

Kirwan's Mineralogy,

when rubbed. This smell is known by the name of Chip. IV. he bituminous odour.

- 4. They become electric by friction, though not inplated t.
- 5. They melt when heated, and burn with a strong mell, a bright flame, and much smoke.
- g. They are insoluble in water and alcohol, but dissolve most commonly in ether, and in the fixed and volatile oils.
- 7. They do not dissolve in alkaline leys, nor form map.
- 8. Acids have little action on them; the sulphuric carcely any: the nitric, by long and repeated digestion, dissolves them, and converts them into a yellow substance, soluble both in water and alcohol, and simihar to the product formed by the action of nitric acid on

The bitumens at present known may be reduced to three; namely, asphaltum, mineral tar, and mineral caoutebouc. Bitumen has been found also united to R restnous compound, in a curious substance first accurately examined by Mr Hatchett, to which he has given the name of retinasphaltum. United to charcoal in various proportious, it constitutes the numerous varieties of pit-roal, so much employed in this country as fuel.

1. Asphaltum. This substance occurs in great a. Asphaltum. bundance in different countries, especially in the island of Trinidad, on the shores of the Dead Sea, and in Albania, where it is found in vast strata. It is supposed

Book II. Division II. that it was first liquid, and that it acquired solidity by exposure to the air.

Its colour is black, with a shade of brown, red, or grey. Its specific gravity varies. That of Aban-a, as ascertained by Klaproth, was 1.205°; but it was somewhat contaminated with earth. Kirwan, in parer specimens, found the specific gravity to vary from 1.07 to 1.165†. Klaproth has lately published an analysm of the asplialtum of Albania.

Properties.

He found it insoluble both in acids and alkalies, as also in water and alcohol; but soluble in oils, petroleum, and sulphuric ether. Five parts of rectified petroleum dissolved one part of asphaltum without the assistance of heat, and formed a blackish brown solution, which by gentle evaporation left the asphaltum in the state of a black brown shining varnish. The solution in ether was of a pale brown red colour; and when evaporated, the asphaltum remained in the state of a semi-fluid substance of a reddish colour, still insoluble in alcohol.

Action of beat.

A bundred grains of this asphaltum being distilled in a retort, by a heat gradually raised to reduces, yielded the following products:

100

· Beltroge, W. 385.

These ashes consisted chiefly of silica and alumina, with one iron, lime, and manganese *.

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The asphaltum found in Albania is supposed to have Uses. instituted the chief ingredient of the Greek fire. The By ptians are said to have employed this bitumen in balming. It was called mumia mineralist. The antents inform us that it was used instead of mortar in sailding the walls of Babylon.

2. Asphaltum is seldom absolutely pure; for when sobol is digested on it, the colour of the liquid beomes yellow, and by gentle evaporation a portion of etroleum is separated 1. Mineral tar seems to be Mineralter, othing else than asphaltum containing a still greater roportion of petroleum. When alcohol is digested on a considerable quantity of that oil is taken up; but here remains a black fluid substance like melted pitch. ent acted upon by alcohol, and which therefore appears' possess the properties of asphaltum, with the excepion of not being solid . By exposure to the air, it is mid to assume gradually the state of asphaltum.

3. Mineral caoutchouc is a singular substance, hither. Mineral cafound only in Derbyshire. It is soft and very clasie, not unlike common caoutchouc, or Indian rubber, and, like that substance, it may be employed to efface pen-

Raproth's Bettroge, in. 316. + Watson's Chem. Ermye, iti. 4. 1 Hatchett's Observations on the Change of some of the Principles of Vere-See into Bitumen. Phil. Trans 1804.

Chemists and mineralogues have united mineral far to petroleum a variety If it be true that pure petroleum is insoluble in alcohol, arrangement is right; but I suspect a mistake. I had no opportuof trying supbibe, but the purest petroleum I have been able to re readily yields to alcohol, unless it has been left exposed to the air.

B L II Divocen II. cil marks from paper; but it soils the paper a link. Its colour is dark brown, sometimes with a share; preen, at other times of red. The first account of it was published by Dr Lister in the Philosophical Transctions for 1673. It occured in an old forsakea moce. Derbyshire. He calls it a subterraneous fungus, man uncertain whether it belongs to the vegetable or mineral kingdoms; but rather inclines to the former opinion, and hints that it may have grown out of the od birch props used in the mine. It was first accuracly described by Mr Hatchett. Delametheric first entitled its properties; and an analysis of it has been lably published by Mr Klaproth.

Properties.

According to Klaproth, it resists the action of almost all liquid menstrua; neither alcohol, alkalies, nor name acid affecting it. Even oils were not found by him to dissolve it, though Delametherre assures us that he obtained a solution of it in olive oil? Petroleum succeeded best with Klaproth, assuming a bright yellow colour, while the caoutchouc was rendered transparent? When heated it melts, takes fire, and burns with a bright flame and a bituminous smell. The melted mass still continues adhesive, and may be drawn out into threads. It is now soluble in oils §.

Action of heat. One hundred grains of this substance being distilled in a retort gradually raised to a red heat, yielded the following products:

⁴ Vol. wii. p. 6179.

Beirege, in. 109.

Jur. de Phys. 222i. 312

[|] Kiaproth, Ibid.

	Grains.	Chap. IV
38 cubic inches (German) of heavy inflam-		
mable air	13.75	
mable air		
A brown thin bituminous oil		
Water slightly acidulous	1.5	
Charcoal	6.25	
Ashes	5.2	
-		
1	00.00	

The ashes consisted of lime and silica, with some iron. sulphate of lime, and alumina .

Mr Hatchett supposes the elasticity of this substance to be owing to a quantity of air confined between its pores.

4. The retinasphaltum of Mr Hatchett is a substance Retinahitherto found only accompanying Bovey coal, in Devonshire. It was first mentioned by Dr Milles; but we are indebted to Mr Hatchett for every thing known respecting its chemical properties.

This substance has a pale brown ochre yellow colour. It is very brittle, and breaks with a vitreous fracture. Its specific gravity is 1.135. When held in the hand for some time, it emits a slightly resinous smell. When heated it melts, smokes, burns with a bright flame, and emits a fragrant odour, at last tainted with a bituminous smell. The melted mass, when cold, is black and brittle, and breaks with a glassy fracture.

Water does not act upon it; but it is partially dissol-

Spinalt int.

^{*} Klaptoth, Beitroge, iii. p. 119.

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ved by alcohol, potash, and nitric acid; the dissolved portion having the properties of a resin; the undissolved, of asphaltum. Mr Hatchett analysed it, and found it composed of *.

55 resin

41 asphaltum

3 earths

99

Pit-coal, of three kinds.

5. Pitcoal, one of the most useful of all the mineral productions, has been divided by mineralogists into various species, according to its external appearances, and the nature of the strata in which it is found; but in a chemical point of view, its most important varieties may be distinguished into three sets: 1. Those that still contain several vegetable principles, strictly so called, and thus give evident marks of their origin. To this head belong most of the varieties of coal arranged by Werner under the head of brown coal. Mr Hatchett has shown, that in some of the substances belonging to this class there is to be found a portion of vegetable extract; in others, as in Bovey coal, a portion of resin; besides the charcoal and bitumen, which constitute the greatest part of its constituents. another species of brown coal Klaproth obtained, by means of alcohol, a brown red tincture, which left behind it on evaporation a reddish bitter extract partially soluble in water +. 2. Those kinds of coal that contain no traces of unaltered vegetable principles, but

coal,

E. Brown

2. Black coal,

^{*} Hatchett, on the Change of some of the Principles of Vegetables to Bitumen. Phil. Trans. 1804.

[†] Beitrage, iii. 322.

we composed of various proportions of bitumen and Chap. IV. harcoal, contaminated like the former with earthy natter. To this head belong the varieties of coal aranged by Werner under the name of black roal, which bound so much in Britain. Kirwan has given us a ery ingenious analysis of several specimens, founded upon the property of nitre to deflagrate with charcoal, ut not with bitumen. By throwing coal reduced to mall pieces into a given weight of melted nitre, he obained a deflagration, and judged of the proportion of harcoal present by the quantity of nitre decomposed *. The following is the result of his analyses.

100 Parts.	Charcoal.	('amen.	Earth.	Sp. Grav.
Connel coal	75.2	21.7	3.1	1.232
Slate coal	47.6	32.5	19.9	1.426
Whitehaven	57.0	41.3	1.7	1.257
Wigan	61.7	36.7	1.0	1 268
Swansey	73.5	23.1	3.4	1.357
Leitrim	71.4	23.4	5.2	1.351
Newcastle	58	40	2	1.271

A number of experiments on the constituents of diferent specimens of British coal have been lately ublished by Mr Mushet †. He subjected the coals to istillation in close vessels, and thus ascertained the poron of volatile matter which they contained. By burnig the coals he determined the portion of earthy matter reach. The following Table exhibits the result of his rperiments.

^{*} Kirwan's M nerology, ii. 14.

[†] Phil Mag. xxxii. 140.

Book II. Diveina II.

	Volatile Matter	Charecal.	Ashes.	of the coal	Sp. Grs. .f the cake
Welsh furnace coal	8.2	85.068	3.43_	1-337	1
Alfreton furnace coal	45.5	521456	21044	11235	less than
Butterly furnace coal	42 83	521862	41284	1:264	
Welsh stone coal	8:00	891760			
Welsh slaty coal	9:10	84.175	61725	_	
Derbyshire cannel coal	47100	48.307	4.635	11275	
Kilkenny coal	4.25	92.877	21873	1.005	110508
Stone coal under basult	16166	100-740	13'000		
Kilkenny slaty coal	13:00	80:475	61525	11446	
Scotch cannel coal	\$6.51	391430	41600		
Boolavooneen coal	_	82 960	31240	1.435	1-596
Corgee do Frish	9.10	87.491			
Queen's county do.)	10:30		3.1 10	1:403	1.0218
Stonewood, Grant's Caus.					1
Oak wood	20.00	19:500	0.200		

3 Glanos

of unaltered vegetable principles nor of bitumen, but seem to consist entirely of charcoal contaminated with some earthy matter. To this head belong the different varieties of coal arranged by Werner under the name of glance-coal. This kind of coal is less common than the preceding. It is remarkable for its metallic lustre, and for the slowness with which it consumes. Many specimens of this coal have been subjected to chemical analysis; the result was always the same. I shall subjoin a few specimens.

The first analysis by Kirwan (Alineralogy, n. 528.) The others published by Hericard de Thury. See Genlen's Jose. v. 323.

100 Parts	Charcoal.	Earth.	Chap. IV.
Kilkenny coal	97.3	3.7	بشكشب
Anthracite	90.0	10.0	
Ditto	72	20	
Ditto	97:25	2.75	
Coal of Notre Dame de Vaux	x 78·5	20	

When coal is distilled, the products vary according to Producta the class to which the mineral belongs. The coals of the third class yield no volatile products whatever. Those of the second give out abundance of heavy inflammable air; a bituminous oil, at first fluid, then of the consistence of tar; and water impregnated with ammonia. The residue is coke, a species of charcoal applied with the greatest advantage to many of our ma-It burns long and equally, and yields a nufactures. great deal of heat. The oil has been applied by Lord Dundonald to the various purposes of varnish. Coals belonging to the first class yield abundance of heavy inflammable air, a bituminous oil, and water either very slightly impregnated with ammonia, or containing a quantity of acetic acid, as has been ascertained by the experiments of Klaproth and Hatchett.

from coal

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CHAP. V.

REMARKS ON THE PRIMARY COMPOUNDS.

Such are the properties of the Primary Conforms a class of bodies which has been longer known than my other, which has been examined with peculiar care, and which comprehends many of the most important instruments of chemical investigation.

Division.

The substances included under the name of primay compounds were divided into four sets; namely, oxide, acids, colorific acids, and compound combustibles. The two first sets are counterparts to each other; both are composed of the same ingredients, and both admit of the same subdivisions. They consist of oxygen united to the simple combustibles, incombustibles, or metals, and are distinguished by their acid properties, or the want of that characteristic. They may be divided map products of combustion, supporters of combustion, and combustibles.

Products of combustion.

Every known product of combustion is a primary compound; for it is remarkable, that in all cases of combustion, the bodies which undergo it, however complicated at first, always arrange themselves so as to form the simplest possible combinations. The apparent exceptions consist of substances which have not in reality

indergone combustion. All the products of combust in known at present are the following:

Chap. V.

- 1. Water;
- 2. Carbonic scid :
- 3. Acid of phosphorus;
- 4. Acid of sulphur ;
- 5. Metallic oxides.

he like manner all the impus emporters of combustion Supported. clong to this class, and are of course primary communds, excepting oxygen itself. These substances are decomposed by the action of heat; a property which satinguishes them very readily from the products. The allowing are all the supporters known to exist, exceptby buygen.

- 1. All the compounds of axote with oxygen.
- 2. All the compounds of muriatic said with oxygen.
- 3. The metallic acids.

The nature of the combustible oxides and acids has been at imperfectly investigated. Comiderable obscurity will hange over them. As chemical agents, they are eferior in importance to oxide and acid products and apporters, and much more liable to undergo changes in meir constitution. With a few exceptions, they are triple compounds, containing oxygen united to two combustible bases, most commonly hydrogen and car-Sometimes azote seems to enter into their conbon. witution.

The colorific acids, strictly speaking, belong to the Colorife name class, and were separated merely on account of the different purposes to which they are applied in chemical investigations.

The compound combustibles, there is every reason to

Book It.

Compound, combustibles. believe, are more variable in their constituents than the preceding classes, and accordingly are more liable to change the r nature. Chemists have not yet been about to anal se them in a satisfactory manner; but the present state of our knowledge leads us to conclude, that as far as their constituents are concerned, they may be arranged under three classes, namely, 1. Those that we composed of bydrogen and carbon. Ether and the volattle oils, when in a state of purity, seem to belong to this class. 2. Those that are composed of bydroga, carbon, and prigen. This appears to be the case with alcohol, with most of the fixed oils, and with those tolatile oils that have been exposed to the air, or are beginning to lose their fluidity. These oils, by that change, lose their smell and most of their distraguishing qualities; but they are in some measure testored by distillation with other, which seems to bring them back to their first condition. 3. Those that are composed of bydrogen, carbon, oxygen, and anote. This appears to be the case with the bitumens; at least if me are to judge from the oil, water, and ammonia, which they yield when distilled.

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DIVISION III.

OF

SECONDARY COMPOUNDS.

By the term Secondary Compound is meant a combination of salifiable bases or primary compounds with each other. These combinations are numerous. Thus acids combine with alkalies, with earths, and with metallic oxides, and form compounds called salts; the earths combine with the fixed alkalies, and form glass; oils combine with alkalies, and form soaps. These combinations come now under our consideration.

The secondary compounds, as far as we are at present acquainted with them, may be arranged under the five following classes.

1. Combinations of earths with each other, and with metallic oxides.

Arrangement.

- 2. Combinations of earths with alkalies.
- 3. Combinations of acids with alkalies, earths, and metallic oxides.
- 4. Combinations of sulphureted hydrogen with alkalies, earths, and metallic oxides.
- 5. Combinations of oils with alkalies, earths, and metallic oxides.



· El Verreintigener

Chip &

CHAP. I.

COMBINATIONS OF EARTHS.

with each other and with metallic oxides, and geompounds distinguished by various names to their external appearance. Some of these tions are formed into vessels known by the stoneware, paraelain, enamel, &cc. many of e of great importance in manufactures and do-

class of bodies is of the highest importance; to not hitherto been examined by chemists with tee of attention to which it is entitled. A few the compounds into which the earths and the oxides are capable of entering have been formiall fewer of these have been described with the subject indeed is difficult, and till late-

The subject indeed is difficult, and till lateest susceptible of accuracy, because the properest earths, and the methods of obtaining them
tre unknown. It is true, indeed, that the greater of these compounds exist ready formed in
tral kingdom, constituting the solid basis of this
fours. But notwithstanding the rapid progress
the analysis of minerals has lately made, it is
y uncertain whether it would be safe to trust
er the result of these analyses; especially as we

imperfectly known.

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are not able to form artificially compounds similar to those which exist ready formed in the earth.

It will be better, therefore, to refer the consideration of these numerous native compounds to the Second Pin of this Work, and to satisfy ourselves at present with general view of the compounds which the earths at capable of forming, a description of those mixing which are employed in the manufacture of stonewin and porcelain, and an account of those combination of earths and metallic oxides which form the different enamels with which these substances are covered. There topics will form the subject of the two following &c. tions.

SECT. I.

OF EARTHY COMBINATIONS IN GENERAL.

THAT several of the earths have a strong affinity for

each other, and are therefore capable of forming combinations, is a fact now well known to the chemical works Difficulty of But it is by no means an easy matter to form them. combinations artificially. If the earths be mixed together in a state of powder, they do not combine atimately; and as few of them are soluble in water, recourse cannot be had, except in a very few cases, to the intervention of that liquid. Chemists, therefore, when they wish to combine the earths together, have scarcely any other agent to employ except fire. This agent answers exceedingly well in combining the metals with

each other, and converting them into alloys: But with

combining earths.

"Espect' to the earths' the case is very different; for no thre which can be produced is sufficiently violent to melt considerable quantity of any of the earths, barytes and strontian excepted.

Chap. L.

It is true indeed that silica, and perhaps all the other Their fuearths, may be melted by a process invented by Saussure. He cements a very minute particle of quartz, or any other substance whose fusibility is to be tried, to a slender fibre of cyanite*, and exposes it in that state to the action of the blow-pipe. He supposes the intend sity of the heat inversely proportional to the diameter of the globule produced. Any heat therefore whatever resay be communicated by means of the blow-pipe, by diminishing sufficiently the bulk of the particle subjects ed to its action. . The limit of intensity is when the particle cannot be farther diminished without rendering the globule produced too small to be seen distinctly by a microscope, and to be measured by means of a micrometer. Saussure found that the heat necessary to melt quartz is equal to 4043° Wedgewood; that which melts alumina is 18900° Wedgewood+:

sing tem-

The temperature, then, at which the fatths mielt is Loweredby inconceivably high: but the quantity of these bodies which can be in this manner subjected to experiment, is by far too minute to draw much satisfactory information from it respecting the nature of the compound which the earths are capable of forming'with each'other. Luckily it is in many cases not necessary to have recourse to it. It is well known that several of the me-

This is a blue colouted trainparent stone, to be afebrwards described.

^{*} Jour. de Phys. 21v. 3.

Book II. Division III.

tals, which require a very high temperature to him them to fusion, melt readily when mixed with a portion of some other metal. Thus platinum, the most infusible of them all, melts readily when combined with ersenic. The same thing happens with several of the carths, as Kunkel first discovered: alumina, for instance, though perhaps the most refractory of them! all, enters very speedily into fusion when mixed with a proper proportion of silica and lime. Chemists have taken advantage of this property. They have might the earths together in various proportions, and then subjected them to hest. The affinity between them was judged of by the degree of fusion which they under went. Mr Pott was one of the first chemists who laid open this method of proceeding. His Lithegrogueria, which was published in 1746, must have been the result of immense labour. It may be considered as the harbinger of all the mineralogical discoveries which fold lowed. Achard of Berlin published, in 2780, a long list of experiments, in which he exposed various miztures of earth to the heat of a porcelain furnace. This was the first direct set of experiments on the combins. tions of earths; for Pott, and Macquer and Darcet who followed him, had confined themselves to native combined nations. Mr Kirwan, in 1704, published a still more accurate set of experiments + on the same subject. Since that time the combination of the earths has occupied the attention of Morveau ; not to mention the interesting experiments of Klaproth | and Saussure ||

Jours de Phys. Bland.

Ellevery.

[•] Mem. Berlin, 1780, p 69.

⁺ Minor slogy, i. 49.

I Jour. de l'Arole Polytacha, I. iii. 198, and Ann. de Chino. 222i. 216.

⁽ Klaproth's Beliroge, i. t.

and the ingenious speculations of Saussure and Dolo- Chap. I. mieu . But the most important experiments on this subject are those of Darracq + and Chenevix 1; because they have been made with substances in a state of purity, and with the proper precautions to prevent erroneous results. From the experiments of these philosophers, and from the accurate mineralogical analyses of Klaproth and Vauquelin, the following consequences may be drawn.

- 1. There is an affinity between most of the different Action of earths: But this affinity varies considerably in intensi- on each ty. Certain earths combine readily with each other in almost every situation, while others cannot be combined without difficulty. There are some combinations of earths which are found native and abundant; others. on the contrary, seldom or dever occur.
- 2. Certain earths when mixed together become very fusible, while other mixtures are as refractory as the simple earths themselves. The affinity between the earths must not be judged of by this increase of fusibility; for several earths which have a strong affinity for each other do not form fusible mixtures at all. In general, mixtures of earths are fasible only when mixed in certain determinate proportions.
- 3. The three alkaline earths, lime, barytes, and stron- Alkaline tian, resemble one another in their disposition to unite with the other earths. Like the alkalies, they combine with alumina and silica, but show no affinity for magnesia nor for each other. Their action on the new earths has not been examined.

Jour. de Phys. slv. 3. passem.

[†] Ann. de Chim. al. 52.

[#] Phil. Trans. 1803.

Book fl. Division fll. r barytes.

When barytes and alumina are boiled together ma sufficient quantity of water, they combine, and from two compounds - one of which, containing an except barytes, remains in solution; the other, containing a excess of alumina, is in the state of an insoluble por-When barytes and silica are fused together in t platinum crucible, they combine, and form a frank blistered mass, of an apple green colour, which a ge luble in all the acids, but imperfectly soluble in with If this compound be boiled in water, it separates un two portions: one, containing an excess of baryies, dasolves, but the presence of the silica prevents the burner from crystallizing; the other remains in the state of an insoluble powder. Barytes, purified in the usual way, always contains a portion of silica, which it probably takes from the crucible in which it is prepared. So great is the affinity of barytes for silica, that it separates silica from potash t. The affinity between also mina and barytes is not strong enough to separate these two earths when the solutions of them in the same said are mixed together. Muriates of barytes and alumina. for instance, when mixed together, afford no precipitate. provided the salts be pure 1.

Barytes is usually found native combined with acide; but it occurs sometimes united to silica; and the state inlite, which is a transparent crystallized stone, is a compound of silica, alumina, and barytes.

2 Strong-

4. Strontian resembles barytes exactly in its affinition

^{*} Vanquel'n, Ann de Chim axiv 273. † Morveau, thid. 221. 14. † Darraeq, thirt al. 57 — Cheneviz un German, p. 17.—Phil Tom. 1802.

for the earths. It unites precisely in the same way Chap. I. with alumina and silica, and precipitates silica from potash. It shows no tendency to unite with magnesia; for is its affinity for alumina strong enough to occasion: precipitate when the muriates of strontian and alumiare mixed together. It shows no tendency to uniter with barytes.

Strontian always occurs native combined with acids; least it has never yet been found united to earths.

5. Like the other alkaline earths, lime has a strong 3. Lime Minity for alumina and silica. Scheele first observed. that when alumina is mixed with lime-water, it combines with the lime, and leaves the water in a state of purity . The compound thus formed is insoluble in water. Chenevix has shown, that this affinity between lime and alumina facilitates the solution of lime in a fixed alkali. When a solution of potash is boiled apon pure lime, no more of it is dissolved than would have been taken up by the water alone which holds the potash in solution; but when potash is boiled in a mixture of lime and alumina, this last earth is dissolved, together with a much greater proportion of the lime than can be ascribed to the action of the water alone t. The affinity, however, of these two earths for each other is not strong enough to occasion a precipitate when their solutions in the same acid are mixed together. Thus no precipitate falls when muriates of lime and alumina are mixed t.

Gadolin first remarked that lime-water forms a pre-

5 1 12 9 1

[†] Phil. Trans. 1802, p. 346 4 Schecle, i. 196

Darracq, Ann. de Char. 21, 58, and Chenevis, Ibid.



A comment to the comment of the comm

Chap. 7.

be present, the alkaline carbonate throws down of the earths in combination. When potash is upon this compound of magnesia and alumina, wery small proportion of the alumina is dissolute greater part being retained by its affinity for ignesia. By dissolving the residuam in muriatic and precipitating by carbonate of potash, a portif the magnesia is retained in solution. Potash on the residue dissolves a new dose of alumina. Thus we see that the separated from each other. Thus we see that then of magnesia on alumina is the reverse of the of alumina on lime; the last promotes the solution alumina.

gnesia, when mixed with silica, forms a fusible and; but the mixture of this earth with barytes, trontian, with lime, or with alamine, is not fusithe heats of our furnaces.

ber earths. These native combined with ber earths. These native combinations consist mes of two earths, sometimes of three, and in getime metallic oxide is also present. They may spechended under three classes:

- 1. Alumina and magnesia,
- 2. Silica and magnesia,
- 3. Silica, alumina, and magnesia.

steatites and kiffekille; to the third, the eyanite

Chenevis on Corunden, p. 67.

Book II. Division III. characteristic of the earthy combinations which combat magnesia; but the ruby and cyanite are both of them transparent and hard. It is worthy of attention, the magnesia enters into fewer fusible compounds than any of the alkaline earths.

< Alemina,

7. Alumina, as has been already remarked, ba an affinity for all the alkaline earths. It has also in affinity for silica. When silicated and aluminated potash are mixed together, the mixure in about an hour becomes opaque and gelatinous, evidently in consequence of the combination of the two earths *. When this precipitate is dried and analysed, it yields both the lica and alumina. The presence of alumina, as Mr Chenevix has remarked, facilitates the solution of siles. in potash. When a mineral containing both silica and alumina is fused in the usual way with a sufficient quantity of potash, and then dissolved in muriatic seid. a number of white flakes frequently remain, which the acid is incapable of dissolving. These white flakes are pure siliceous earth. Their quantity is greatest when the proportion of the silica over the alumina is greatest When the proportion of alumina is considerable, the insoluble residue is less, and in some cases at disappear altogether +.

Alumina enters into fusion with none of the other earths except lime, and not even with the fixed alkalies. In this respect it differs exceedingly from silica; which from the numerous fusible compounds that it is capable of forming, was formerly called the vitrifiable earth.

^{*} Morvezo, Ann de Climarra, 848. | | that, 12viii. 201

But alumina enters into several triple earthy mixtures which are fusible. Thus mixtures of alumina and silica with any of the alkaline earths, and with magnesia, are fusible.

Chap. I.

Alumina is found native in great abundance, and almost always combined with other earths or with metailic oxides, for which it has a very strong affinity. The native earthy combinations, of which it constitutes an ingredient, may be divided into six classes. These are,

- 1. Alumina and magnesia,
- 2. Alumina and silica,
- 3. Alumina, silica, and barytes,
- 4. Alumina, silica, and lime,
- 5. Alumina, silica, and magnesia,
- 6. Alumina, silica, and glucina.

To the first of these classes belongs the ruby; to the second, mica, chalcedony, &cc.; to the third, staurolite; to the fourth, zeolites; to the fifth, cyanite; and to the sixth, the emerald. These minerals will be described hereafter. All the compounds into which a considerable portion of alumina enters are pretty heavy, and several of them are exceedingly hard.

S. Silica, as we have seen, has an affinity for the al- 6. Silica. kaline earths and alumina. It has likewise an affinity for zirconia. When a mixture of this last earth and silica is heated strongly, it shows a tendency to fusion . It fuses, as we have seen already, with barytes, strontian, lime, and magnesia; so that alumina is the only earth hitherto tried with which it does not enter into fusion.

^{*} Morveau, Ann. de Chim. RETi. 259.

Book U. Division III. Silies is found native more abundantly than any of the other earths. It also enters into a greater vancy of combinations, but almost always with earthy bodies. The combinations of which it is known to constitute a part may be reduced under eight classes; namely,

- 1. Silica and alumina,
- 2. Silica and lime.
- 3. Silica and magnesia,
- 4. Silica and zirconia.
- 5. Silica, alumina, and lime,
- 6. Silice, alumine, and magnesia,
- 7. Silica, alumina, and glucina,
- 8. Silice, alumina, and barytes.

All of these classes have been mentioned already, and instances given of minerals belonging to each, except the fourth, to which belong only the wircon and the by acintb.

The most beautiful earthy combinations are those which consist chiefly of silica and alumina. To the class belong almost all the gems. The combinations of these two earths are also the most useful; for they constitute the basis of porcelain, stoneware, glass pots, and all the different earthen utensils.

Combinations of metallic oxider and earths. 9. Several of the earths are capable of combining likewise with metallic oxides, and of forming compounds; the nature of which has scarcely been examined by chemists. All the oxides which fuse readily into a glass are capable, in that state, of combining with earths by the assistance of heat, and of forming opaque compounds, known by the name of example; some of which will come under our consideration afterwards.

Hitherto only six metals in the state of oxides have en found native combined with earths. These are,

Chape to

- 1. Chromium.
- 4. Zinc.
- 2. Nickel.
- 5. Manganese,
- 3. Copper,
- 6. Iron.

Chromium constitutes very frequently the colouring 1. Chromimatter of the ruby, in which it is combined with alumina and magnesia; but this metal does not appear to have any remarkable tendency to combine with earths except when it is in the state of an acid.

Nickel has been detected by Klaproth in the chryso- s. Nickel. mase, and in the green matter called pimelite which cometimes accompanies it; but whether the combinason is chemical, or merely a mixture, has not been aspertained.

The oxide of copper shows no peculiar tendency to 3. Copper. mite with earths, and hitherto it has been detected in me earthy compound only; namely, the smaragdite, a mineral which contains 1.5 per cent. of oxide of copper. But whether this oxide be chemically combined or merely mixed with the other ingredients is not known.

The oxide of zinc has been found native combined 4. Zinc. with silica in different varieties of calamine. these bodies are chemically combined has been demonarated by Mr Smithson in his valuable paper on calamines .

It has been found likewise combined with alumina in a singular mineral lately analysed by Ekeberg, to which be has given the name of automalite. This mineral is of a dark green colour, crystalkzed in octahedrons Brok 1. Dorina III. composed of this passed. Where and the course crystals are appeared it makes quarte, it makes quarte, its specific growth that were not are powder has a light green smane. Hence he is it does not men nor entage me smane. Hence he is into a gione, common where his makes into a gione, common where his makes into the spinel, was found common of the makes and 244 vaide of zinc; & common of the makes and 244 vaide of zinc; & common of the makes and 244 vaide of zinc; & common of the makes and 244 vaide of zinc; & common of the makes and 244 vaide of zinc; & common of the makes and 244 vaide of zinc; & common of the makes and 244 vaide of zinc; & common of the makes and zinc and zinc

g. Metrger noce. But it is in the oxides of management and in cially the latter, that the affinity members are a metallic oxides is most conspicuous. The manyanese occurs paire combined with implication of the binese and in loured stones, as school, garnet, i.e. But with ned with earths, if we except buryers, a improved with earths, if we except buryers, a improved the litherto observed, a very small proposition compound.

6. Iron.

The oxide of iron is one of the most committee to minerals. It is found in a scare of tion with almost every earth. To it indeed cases, is the colour of minerals to be ascribles than seven distinct colours, besides a gre of shides, are observed in minerals contains and to the iron in most cases is the colour to bed. These colours are,

- 1. White......The sommite, &cc.
- 2. Black.....Obsidian

Gchlen's Jour. v. 442.

3. Green.....Euclase, thallite

Chap. t.

- 4. Blue.....Lazulite, native prussian blue
- 5. RedGarnet, ruby
- 6. Y. ilow Topaz
- 7. Brown......Tourmaline

com the experiments of Kirwan, we learn that forms a fusible mass when mixed with barytes, alumina, and silica, when it exceeds the proporof earth considerably. With magnesia it forms an mel, but scarcely fuses completely. It renders a ture of alumina and silica fusible at a very low heat in the proper proportion.

D. During the fusion of several mixtures or combi- Earthy as of earths, especially those which are found nathe mixture very often frothes considerably, and foth au-It a great number of air bubbles. The emission of air hubbles can only be referred to one or other of causes. Either a quantity of water leaves the mare in the form of steam, or the metallic oxides, th are almost always present, are reduced, and alaheir oxygen to escape in the form of gas; or, the earths themselves undergo some change, Blet go a portion of their oxygen in the form as. It is probable that the phenomenon is partiwing to all these causes; for these earthy mixtures much lighter after having been exposed to the acof the fire than they were before. Thus a mixof equal parts of silica and magnesia, after being led to 134° Wedgewood, lost 0.195 of its weight; a mixture of equal parts of alumina and barytes

Book II. Division III. lost in the same situation 0.275 of its weight. Not so great a loss as this cannot be ascribed to my the cause than the evaporation of water.

Again, several minerals which are coloured by the tallic oxide become transparent when exposed to the violent heat; a change which must be occasioned to the revival or dissipation of the metal. This is at ease, for instance, with the brown obsidian lava of D. lomieu, with the lazulite, and with school? Now is exceedingly difficult to free the earth completely from all metallic bodies; especially alumina, the earth not remarkable for giving the property of frothing to earth mixtures.

Concerning the third cause of the frothing of mine rals, namely, the decomposition of the earths then selves, nothing like precision could be expected all the component parts of the earths were discovered. Herebolds announced some time ago, that the earths have the property of absorbing oxygen from the atmosphere. In that case the frothing might, in some instances at less, be ascribed to the emission of this oxygen on the spotcation of heat; but the experiments of Humboldt have been tried without success by Saussure, Fabroni, Champy, Berthollet, and Delametherie 1. He has lately, however, announced that he has repeated his experiments successfully in concert with Gay-Lassac; he even declares his belief that earths are capable of combustion as well as other bodies f. Mr Humboldt his not yet published a detailed account of these latter ea-

[•] Morveau, Jour de l'Ecole Polytethique, L iii. 306.

Saveure, Jour. de Phys. xlv. 16.

[#] Ibid. Iri, 6a.

[€] Gehign's Jour. v. 232,

meriments; of course, we are ignorant of the way in which they were performed, and of the degree of probability which they give to his opinion: an opinion concerning which it would be unfair to decide without farther evidence.

Chap. L

We now know from the discoveries of Mr Davy, that the bases of the alkaline earths are very combustible metals. These earths of course are products of combustion, and therefore incombustible. If Mr Humboldt's opinion referred to the bases of these earthy bodies, it was correct; but if it referred to the earths themselves, it was erroneous.

11. There is another phenomenon respecting the fu- Sometimes sion of earthy combinations which has not been hither- ble fusion. to explained. Some minerals, when exposed to the action of heat, melt very readily, and generally with a good deal of frothing; but if the heat be continued, they soon become solid again, and do not fuse unless the temperature be considerably augmented. This is the case with prebmite, thallite, and some felspars. Thus the prehmite undergoes its first fusion at 21° Wedgewood, its second not till it be heated to \$100. It cannot be doubted that in this case some one or other of the ingredients which contributed to the first fusion must be driven off; especially as the mineral after fusion loses entirely its original appearance and properties. The late important experiments of Sir James Hall on the exposure of carbonate of lime to strong heats, under such a pressure as prevents the escape of the carbonic acid, have shown that this substance, under these

Sansoure, Jour. de Phys. 214. 24.

Book II. Division III. the fusion of the carbonate of lime, which he has a complished in a great variety of instances. It was not be surprising if the first fusion of the zealute are owing to the presence of carbonic acid, and the fractal to the escape of that substance. This conference sees to have struck Sir James Hall, whose skill in that the cult department of chemical investigation will no doubt enable him in a short time to confirm or refute it.

Often depends on evolung.

12. A change no less curious is produced on the appearance and fusibility of some minerals by the size. ness or rapidity with which they are cooled after being subjected to fusion. Whinstone (greenstone of Wesner), for instance, is a mineral which fuses when rand to a temperature between 40" and 55° Wedgewoods If it be allowed after fusion to cool rapidly, it is cose verted into a dark-colouted glass much more faulte than the original whinstone; but if it be made to cool very slowly, it assumes an appearance which has some resemblance to the original whinstone. In this state if is much less fusible than the glass. Sir James Hall, the discoverer of this curious fact, has given the whin in this last state the name of crystallite, a term suggested by Dr Hope. Thus the rock on which Edinburgh Castle is built fuses at the temperature of 45° Wedgewood. By rapid cooling it is converted into a glass, which melts at 22°; by slow cooling into a crystallity! which melts at 35°. The whin of Salisbury Craig fuses at 55°; its glass at 24°; its crystallite at 38° . Lavas agree with these minerals in this respect. The

. Bien. Teber V. 78.



wing Table exhibits the degree of Wedgewood's meter at which the rocks tried by Sir James Hall rgo fusion in their natural state, in the state of glass, in the state of crystallite.

Chap. I.

Substances.	Original State.	Glass.	Crystal- lite.
in of Bell's Mills quarry, greenstone),	40°	15°	320
in of Edinburgh Castle, (por- hyry slate),	45	· 22	35
umns, Arthur Seat, (gr. stone), in near Duddingston Loch,	55	18	85
greenstone),	43	24	38
in of Salisbury Craigs, (ditto), in from Water of Leith,	55	24	38
ditto),	55	16	37
in of Staffa, (porphyry slate),	. 38	141	35
ra of Catania,	33	18	38
nonte,	32	18	36
ra of La Motta,	36	18	36
ra of Iceland,	35	15	43
ra of Torre del Greco,	40	18	28
ra of Vesuvius, 1785,	18	18	35

of these bodies depends upon the length of time are allowed to remain in a higher temperature that in which the vitreous fusion takes place. A ute or two is sufficient to induce the change; but it fore complete the higher the temperature is above point of vitreous fusion while the crystals are form-

^{*} Edin. Trans. v. 75.

Blook II. Dismontfl. ing; provided always it be not so high as to private the crystals from shooting.

The striking difference in the fusibility of these by dies seems to depend upon the way in which the con. ponent parts of the mineral are combined. When even ed rapidly, these component parts have not time to tree bine according to their affinities, but are huddled ungularly together. Of course the force of the affinites of those component parts which have a tendency to ton. bine, must conspire with the action of the fire to destrer the cohesion of the parts; the consequence of what enust be fusion. Whereas in the crystallite, the conponent parts having had time to combine according to their affinities, instead of conspiring with the dwellest action of the fire, oppose it by coinciding with the form of cohesion. In the first case two forces combine to produce fusion; in the second, only one. Something similar to this may perhaps take place in those minerals which undergo a double fusion, the first in a lower heat than the second ..

Opon this subject the reader may consult a curious paper by the line. Mr Gregory Watt, whose premature death must be deeply regressed by the cultivators of chemical science. He fused about seven hundres weight of greenstone in a furnace, allowed it to cool alowly, an , then examine the various changes it had undergone. See Phel. Trans. 1804.

Chap. L.

SECT. II.

OF STONEWARE.

a distinguishing character of alumina is to contract bulk, and to acquire the hardness and solidity of a body, if it be made up into a paste with water, ad slowly in the open air, and then exposed to the og heat of a furnace. It communicates this proper- Properties to the numerous tribe of clays, which are so comin most countries; a property which has doubtsuggested the applying of clays to form various sels and other bodies for the purposes of the arts and domestic economy. Clays consist essentially of aluand silica mixed in various proportions; and baked of course, must be considered as a compound of etwo earths. We shall, in the present Section, take bort view of the manufacture of the various substanformed of baked clay. As there is no single term our language which comprehends all these bodies, I we given to the Section the title of stoneware, which dudes the greater number of them. These bodies be divided into four sets; namely, bricks and tiles, to and eracibles, stoneware and porcelain. A detailed Clay venels count of the manufacture of these bodies would be proper in this place, and is not intended; such a stch only is meant as may lead the reader to the prinles of the art.

Book II. Division III.

I. BRICKS AND TILES.

Bricks are oblong masses of baked clay, used as a substitute for stones in building, and tiles are prepared for covering the roofs of houses. The use of them may be dated from the most remote antiquity; and when properly prepared, they are little inferior in durability to stony bodies themselves. Many ancient monuments built of brick still remain entire.

The day.

Bricks are prepared from the common blue clay, which occurs in abundance in almost all countries. It ought to be of such a nature as not to be liable to fusion when exposed to a strong heat; at least if the bricks are intended to withstand the action of fire. The presence of a portion of lime renders clay fusible. If the clay contains too little sand, the bricks are liable to crack in cooling; too much, on the other hand, prevents the proper degree of cohesion. Hence the goodness of the bricks depends upon the clay of which they are made, which can only be ascertained by experience. It depends also upon the degree of heat employed in baking them. When not sufficiently burned, they are liable to crumble down in a very short time.

Mamilaeture of Bricks The clay is dug out of the earth, and after being exposed for some time to the air is reduced to powder, and formed into a paste with water. The bricks are then formed in moulds, exposed for some time to dry in the open air, and then burnt in a large furnace constructed on purpose. Tiles are formed in the same way. The clay, however, is finer, and it is usually ground in a mill. Bricks and tiles should be impervious to water: They should be capable of withstanding the action

of heat, and not be subject to moulder. The clay of which these substances are made always contains iron; bence the red colour which they acquire in burning.

Chap, L.

II. POTS AND CRUCIBLES.

For various manufactures it is necessary to have vessels capable of resisting very violent heats without fusion, even when in contact with bodies that act the part of fluxes. This is the case for instance in glass making, and in the smelting of the ores of the difficultly fusible metals, as iron and copper. The glasshouse pots are formed of the purest kinds of clay that can be procured. By pure clay is meant a clay free from lime, and from any great proportion of iron. Sturbridge clay is commonly used for the purpose in this country. To prevent it from contracting too much in bulk when heated, it is mixed with a proper proportion of old pots reduced to powder. The pots are baked in a mould with great attention, and afterwards allowed to dry as slowly as possible. They are afterwards baked in the requinite degree of heat.

The crucibles used for chemical purposes are pre- Crucibles. pared of similar materials. Sometimes sand is used instead of old crue bles; semetimes plumbage is substituted. The crucibles are then called black lead crueibles ".

For a detailed account of crucibles and clay fernaces, the reader is referred to Baunte's Chemistry, 116, 213. The best a c ant of furnaver for chemical purposes which has hither to appeared in the English language, in given by Dr. Lewis at the beginning of his Phi - obbical Commerce of the After. Practical chemists will do well to consult that important treatise.

DECEMBER CONFORMER

III. STONEWARE.

The manufacture of versuels of stoneware a simple socient date. It seems to have originated many a Amatic natural, and was probably carried to an East from time immerisorial, pretty much not upon thed in India at present. It was well knows to leave, as we have from the Old Testament, long being the Bobylouish captivity. Successare vessels differ for each other in their fineness, according to the panty of the materials of which they are composed; and send ingly are distinguished in this country by a variety of manes, such as finenesse, prefere ware, queen's war. Wedgewood-ware, bec.

Vessels of stoneware consist essentially of two past.

1. The body of the vessel called the biscuit. 2. The glassy covering with which it is coated called the glate.

The biscuit is composed of two ingredients; the folis a fine white clay, known by the names of robaccapipe clay and petters clay, and the second a fine white sand. In this country the sand is procured by calcining flints to whiteness, and grinding them in a mill. The glaze consists of various ingredients according to circumstances. We shall first take a view of the preparation of the biscuit, and then of the glaze.

Staffordshire is the county in which the greatest quantity of stoneware is made in Britain. Hence it is often called Staffordsbire ware. The clay employed in the manufactures is brought from Dorset and Devonshire, and the flints from the south-eastern counties. The clay is reduced nearly to the consistence of milk

water, and the liquid passed through lawn sieves oally increasing in fineness. By this means all sparse sand is separated, and only the minutest parof the clay left suspended in the water. The flint, tously ground to a very fine powder between millof chert (hornstone), is mixed with water seely, and brought to the consistence of cream. two liquids containing the clay and the flint are together in various proportions according to mstances; the flint liquor, in some cases, amountth, in others to the of the whole. The mixed after being well stirred, is let into troughs, and prated to the proper consistence. This mixture is taken out, and kneaded into a fine dough. This of the process is tedious, but essential; as it is mebry that the whole should be equally mixed, and of same consistence. The dough thus prepared, is for use in damp vaults; from which it is taken as sion requires, and given to the workmen, who it on the lath to the proper shape. The vessels rudely formed are in good weather dried in the air; but in bad weather, in stoves heated for the ose. When sufficiently dry to bear it, they are jed back again to the lath to have their asperities off, and the exact shape given. They are then wed to dry as completely as possible. Afterwards are put into cylindrical earthen vessels, formed of oded fire-bricks and clay, called reggars; and when shape will admit, each seggar is filled with the vesto be baked, laid one above another. These segare piled one above another in a large circular capable of holding a considerable number. When reggars are put in, the door of the kiln is built up

Chap. I.

Book II. Devis on Life with bricks, and plastered over of furnaces built round the ki in the middle of it, are then k kiln and its contents to a very fire is kept up for 48 hours, as The vessels, when taken out biscuit. They resemble a tol and like it absorb water with be employed commodiously, a ty, till their surface is covered

The glaze,

The glaze employed to commy be distinguished into three or sulphuret of lead, common a Galena is employed only to sest kinds of stoneware. Whand exposed to the proper tendissipated, and the lead oxid strong tendency to vitrefy, and earthy bodies. The brown g ware, then, is nothing else than sels are dangerous when used to

s, Common

Common salt is employed to stoneware vessels. The prothe biscuit is sufficiently bake salt is introduced into the k this salt into a vapour, it pene holes left on purpose, and surr atmosphere of salt. The salt surface of the biscuit, disposes

as the glaze is easily corrodec

and various saline bodies.

A Enumel

Enamel is nothing else that over the surface of the biscuit



Chap. L.

various proportions of oxide of lead mixed with ad and glass. Mr Wedgewood published the followig as the enamel which he used : One hundred parts I lead are melted with from 15 to 40 parts of tin, and The mixture oxidized completely, by exposing it to heat an open vessel. One hundred parts of this oxide are pixed with 100 parts of a fine white sand, composed of tree parts silica and one part of tale, and with about parts of common salt. This mixture is melted, then educed to powder, and formed into a liquid of the consistence of cream. According to Dr Watson, the ellow glaze used in Staffordshire is composed of 112 parts of white lead, 24 of ground flint, and six of flint . class, mixed with water to the consistence of cream . The biscuit is dipt into this liquid and drawn out again; by this means its surface is covered with the solid matfer of the enamel, the water being speedily absorbed. The vessels are put into the seggars as before, but a litde cross of stoneware is interposed to keep them sepaparate. They are put again into the kiln and heated as at first. The enamel melts into a glass, and spreads equally on their surface. The excellency of a good enamel is, that it easily fuses into a kind of paste at the heat which is necessary for baking stoneware, and spreads equably on the vessel, forming a smooth glassy surface, without losing its opacity, or flowing complete-By into a glass. Its whiteness depends upon the proportion of the tin, its fusibility upon the lead +.

Chemical Errays, 11. 269.

[†] For an account of the manufacture of stoneware, the reader may consult Baume's Chemistry, iii. 227, and Watson's Chemist Euryr. ii

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the sale of the same of the same of the sale of the sa

open marchine. They were haven to the Language to Common and the deliges of higherdones, or we are advanced by their are the total to the post of the transfer of the post of the

Chap. L.

uring a set of experiments in order to ascertain the est mixtures for making crucibles, stumbled upon a compound which yielded a porcelain similar to the Easten. In consequence of this discovery, Saxony soon coduced porcelain scarcely inferior to that of Japan in eauty, and superior to it in solidity and strength: But a composition was kept secret; nor were there any ecurate ideas respecting the component parts of porcelain among men of science, till Reaumur published his issertations on the subject in 1727 and 1729.

That celebrated philosopher examined the porcelain I Japan and the different imitations of it which had een produced in France and other parts of Europe. The texture of the first was compact and solid, but that I the imitations was porous. When both were expoded to a strong heat, the first remained unaltered, but me others melted into glass. From these experiments a drew the following ingenious conclusions.

Porcelain owes its semitransparency to a kind of semivitrification which it has undergone. Now it may
beceive this two ways: 1. Its component parts may
be such as easily vitrify when sufficiently heated; but
the degree of heat given may be just sufficient to occalon a commencement of vitrification. This pore lain
when strongly heated will easily melt. Such, therefore, was the composition of the European imitations
of porcelain. 2. It may be composed of two ingredients; one of which vitrifies, but the other is not altered by heat. When a porcelain composed of such
materials is baked in a sufficient heat, the fusible part
melts, envelopes the infusible, and forms a semitranspatent substance, which is not farther alreved by the same
degree of heat. Such therefore must be the porcelain

Book II. Division Hi. of Japan. Father Entrecolles, a missionary to China, had sent an account of the Chinese mode of making pos-celain, which coincided exactly with this ingenious thought of Resumur. The ingredients, according to him, are a hard stone called petunse, which they grad to powder, and a white earth called kaolin, which is institutely mixed with it. Resumur found the petunselusible, and the kaolin infusible, when exposed separately to a violent heat.

These notions were not prosecuted farther by Remmur; but in 1758, the Count de Lauragais, assisted by Darcet and Le Gay, began a set of experiments which were continued for four years, and which led to the discovery of a porcelain possessed of the same qualing with that of China, and inferior only in whiteness. Macquer, who at that time superintended the manufactory of Sevres, advised the French government to propose a reward for the discovery of earthy substances capuble of forming a white porcelain. This was done: and in consequee of it, Villaris, an apothecary of Bourdeaux, announced the existence of a white earth new Saint-Yriex la Perche, in the department of the Haute Vienne, which in his opinion would answer the purpose. a It was tried by Macquer with the expected success. A porcelain manufactory was established at Sevres, and at present there are no fewer than 30 in France. Different manufactures of porcelain have been successfully established likewise in England; first at Chelsea in the neighbourhood of London, and afterwards in Coalbrookdale, and in Derby.

[&]quot; Bourmy's Memoire sur les Ouvrages de Terres suites, p. -a.

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The essential ingredient of porcelain is a very pure clay known by the name of porcelain clay. This is equivalent to the kaolin of the Chinese. Macquer and Baumé, in the course of their experiments, ascertained that very fine porcelain may be made by using the earth of alum instead of clay; but the great expence prevents the possibility of introducing it into the manufacture of that article with advantage. Sometimes porcelain clay consists of materials mixed in such proportions that no addition is necessary; the biscuit made from it being susceptible of undergoing that semivitrification which gives the transparency and compact nature which distinguish porcelain. Such is the porcelain clay of Limoges. According to the analysis of Hassenfratz, it is composed of 62 silica

Chap. L.
Por clain
carth.

19 alumina

12 magnesia

7 barytes

100*

20 silica

12 moisture

8 loss

100 +

† Kirwan's Min. i. 179.

Ann. de Chim. 21v. 144.

Book II. Division I'I. Giobert has announced that porcelain earth is sometimes nearly pure magnesia and silica 1.

When an addition is necessary, the substance used is felspar, which is equivalent to the petun e of the Chanese. According to an anonymous German writer, the finest Saxon porcelain is formed by mixing together equal weights of ground felspar and porcelain the The method of forming the biscuit of porcelain is the same as of stoneware, and therefore requires no paracular description.

Glaze of Porcelain. Porcelain is always covered with a glaze, composed of earthy ingredients without any mixture of metalic oxides. Hence the high temperature necessary to fore it, and the property which porcelain vessels have at resisting the action of the most corrosive substances precisely as common glass does. The substance common ly employed is felspar; which is composed essentially of silica and alumina united to some potash, to which the fusibility is to be ascribed. This is the glaze said to be used in Sax ny, and, as Brogniant informs us likewise at the manufactory of Sevres, near Paris.

Method of colouring stoneware.

Vessels both of stoneware and porcelain are commonly painted of various colours. These paintings are often excellent, both in elegance of workmanship and a brilliancy of colours. The colours are given by means of metallic oxides, which are mixed up with other ingredients proper to constitute an enamel, and applied in the usual manner with a pencil.

¹ Nicholson's Jour. zii, 277.

^{\$} Societ de en es accelains, &c. p. 608, at the end of the French translation of Ners's Art of Glass-making,

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On this subject much light has been thrown by the experiments of Wedgewood; and Brogniart has lately published a general account of the processes at Sevres, of which he is director *.

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The process differs a little according to the substance on which the colours are to be applied. When the vessels are covered with enamel, less flux is necessary, because the enamel melts at a low heat, and the colours readily incorporate with it. But this renders them more dilute, and makes it often necessary to retouch them. The colours on enamel generally appear brilliant and soft, and are not liable to scale. The flux is either a glass of flint and lead, or borax mixed with flint glass. The colours are usually made into a paste by means of gum water or volatile oils. Some of them are liable to alteration by the action of the lead on them.

The colours applied upon hard porcelain, or porcelain glazed with felspar, are nearly the same as those applied on enamel, but more flux is necessary. They are not liable to dilution, as the felspar glaze does not melt at the heat requisite for fusing the colours and their flux. They are liable to scale off when repeatedly heated.

Colours are sometimes applied over the whole surface of the porcelain; the flux in that case is felspar. But such colours are not numerous, because few oxides can stand the heat necessary for melting felspar without being altered or volatilized.

Painting on porcelain.

^{*} Pbil. Mog. xiii. 342.

Book II. Division III.

- 1. Purple is given by means of the purple oxide of gold precipitated by the smallest possible quantity of muriate of tin. This oxide is mixed with a proper quatity of powdered glass, borax, and oxide of antimony, and applied with a pencil. It cannot bear a strong heat without losing its colour.
- 2. Red is given by oxide of iron. A migture of two parts of sulphate of iron and one part of alum is calcined slowly, till it acquires a fine red colour when cold. This powder is mixed with the usual flux, and applied with a pencil.
- 3. Yellow is given by the oxide of silver, or by oxides of lead, antimony, and sand; green, by the oxide of copper; blue, by the oxide of cobalt; and violet, by the oxide of manganese.

For farther information on this subject, the reader is referred to the dissertation of Brogniart.

Gilding upon porcelain is performed in the same way as painting. The gold is reduced to the state of an impalpable powder by solution and precipitation. It is mixed up to the proper consistence with oil and a small quantity of flux, and applied with a pencil; the vessels are baked a second time. By this the gold is made to attach itself firmly to the vessel, and by the burnisher it acquires the requisite lustre. Klaproth has lately shown, that platinum may be applied upon porcelain with a similar effect. The fine powder of platinum obtained by precipitating the metal by means of sal ammoniac, and exposing the precipitate to a red heat, is to be ground with a little flux and oil to the proper consistence, and applied with a pencil upon the vessels. By

baking these vessels and burnishing, the platinum acquires the requisite lustre ".

their price too much; but this is avoided by an ingeni- plate print-

Painting common stoneware vessels would enhance Copperous mode of copperplate-printing, said to have been first invented by some person in the neighbourhood of Liverpool. The figure which is to be painted on the vessel is engraven on a copperplate in the usual way, excepting only that it is not reversed as is done in common copperplate-engraving. The paint to be applied to the stoneware is brought to the requisite consistence, put upon the copperplate, and the impression taken off, as usual, upon moist paper, by means of the rolling press. The paper, while still moist, is applied to the stoneware biscuit and pressed upon it. By dipping the biscuit in water and agitating gently, the paper is washed off without injuring the impression upon the vessel, the paint having been made up with oil. The impression upon the paper was reversed, but upon the stoneware it is precisely as it was cut upon the copperplate. The vessel being now baked, the paint is glazed on, and assumes its characteristic colour and brilliancy. By this contrivance any number of vessels may be easily printed with the same figures in a very short time. This ingenious process seems to be at present confined to Britain; at least none of the foreign stoneware that I have had an opportunity of examining, exhibited any marks of having been printed. Neither has any mention of the process been inserted into any work which I have met with.

Nicholson's Jeur. vil. 186.

COMBINATIONS OF EARTHS.

The specific gravity of the different kinds of store.

ware differs considerably according to the compacture of the texture. The following are the results obtains by Dr Watson and Brisson:

	Sp. grav.
East Indian China +	.2.385
East Indian China ‡	.2.346
Limoges porcelain +	. 2.341
Bristol stoneware ‡	.2.340
Flint ware ‡	.2.188
Sevres porcelain †	.2.146
Yellow ware !	.1.988
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4 Giordeol Leonys, il. 252.

† Brisson.

1 Water

CHAP. II.

GLASS.

DILICA, when mixed with the fixed alkalies, and exposed to a strong heat, enters readily into fusion. It melts also when heated along with some of the alkaline earths, especially lime, provided a little alumina be present. These mixtures are very ducule while in fusion, and may be readily moulded into any shape we please. If they be suddenly cooled below the temperature at which they become solid, they retain their transparency, and assume those peculiar properties which belong to the substance called glass. Glass then is a combination of the fixed alkalies or alkaline earths with silica, either alone or conjoined with alumina, brought into complete fusion, and then suddenly congealed. Metallic oxides are sometimes added: they assist the fusion like the alkalies, and communicate frequently a peculiar colour to the vitreous mass.

The method of making glass was known at a very History. early period. According to Pliny, the discovery was owing to an accident. Some merchants, with a shipload of soda from Egypt, had cast anchor at the mouth of the river Belus in Phonicia, and were dressing their dinner on the sand. They made use of large lumps of

Book II. Division III.

soda to support their kettles, and lighted fires use them. The heat melted the soda and the stitteous and together, and the result was glass. For some time alm this accidental discovery the manufacture of gass w confined to the river Belux. This manufacture seem to have been carried to a considerable degree of go fection among the ancients. They mention durke glasses, glass prisms, and coloured glasses of vance kinds. But perfectly transparent glass was consideral as very valuable; for Nero gave L.50,000 for me glass cups with handles; a proof that their process must have been far less perfect than ours. It was un al for them to melt the materials of their glass into black mass called ammonitrum, of which statues we sometimes made. This ammonitrum was again meleand purified by refiners. Glass panes seem to ha been first used in windows in the third century, butthe did not come into common use till long after ".

Nature of

While glass is in fusion, the substances which entitute its composition may be considered as combined with each other so as to form a homogeneous mass a milar to water, holding a variety of salts in solution. If it be cooled down very slowly, the different tendent of the constituents to assume solid forms at peculic temperatures will cause them to separate successive in crystals; just as the salts held in solution in water assume the form of crystals as the liquid is slowly evaporated. But if the glass be quickly cooled down to the point of congelation, the constituents have not

See Dr Falconer's paper on this subject, Manchester Memor, is 95, and Dr Merrets's Preface to his Latin Translation of Nen's Art Glass-making.

Chap. 1L

Time to separate in succession, and the glass remains the me homogeneous compound as while in a state of fuion; just as would happen to a saline solution if sudenly exposed to a cold sufficient to congeal it combletely. Hence it appears that the vitreous quality depends entirely upon the fusibility of the mixture, and be suddenness with which it is cooled down to the point of congelation. The substance, though solid, is precisely the same as to its chemical composition as if were still in fusion; the sunden cooling having fixed the constituents before they had time to assume a new arrangement.

All fusible mixtures of the earths proper with fixed alkalies, alkaline earths, or metallic oxides, may be made at pleasure to assume the form of glass, or the appearance which characterises stone or porcelain, according to the rate of cooling; and glass may be deprived of its vitreous form merely by fusing it, and cooling it down with sufficient slowness to enable the constituents to separate in succession.

Many curious experiments on this subject were Changes in made by Reaumur and Lewis, who pointed out the ingaster fomethod of converting different kinds of glass into an opaque, white, hard, refractory substance like porcelain, which is commonly distinguished by the name of Reaumur's porcelain. Dr Lewis, by a variety of experiments, demonstrated that all kinds of glass could not be converted into porcelain. He succeeded only with those that were composed of a variety of constituents. The reason is obvious; such glasses alone contain ingredients

Phil. Com. of the Arts, p. 230.

that become solid in succession. Green glass mend, ed best with him. Indeed this glass is very aprile to quire a crystallized form. The temperature bestural an the change is that in which the glass is softened with out being melted.

It was the curious experiments of Sir James Haller baselt and greenstone, detailed in the preceding Craster, that first explained upon what the vitreous state substances depends. He found that glass (consisting various earthy bodies) always loses its vitreous se and assumes that of a stone, if more than a minute two clapses while it is cooling down from complete must to the point at which it congests.

There are different kinds of glass in common up this country for various purposes. The finest are sle giarr, of which tooking-glasses are made, and flut-glasses or crystal, used for the finest vessels. These are me feeely transparent and colourless, heavy and brillian They are composed of fixed alkali, pure siliceous and calcined flints, and litharge. The manufacturers conthe proportions of their ingredients with great can

[·] For the failest account of glass-wadsog, the render is referred. treatme on the subject published by Ners, an Italian Dr Merry Logishman, translated it into latin towards the end of the 17th. may, and enriched it with many notes. Kunkel translated the liodrice with Merret i notes into German, and added to the world own transcrees experiments on glass-making, and the w-infection the formulas of Nert Kunkel's work was translated into French, published in a quarto volume in 1753. A very elaborate account gino-making has been published also in the Arts of Meters, design under the direction of the French Academy. More lately, a small a hame on glass-making his been written in French by Loysell. The cours, as he describes them, differ in many respects from those follows in this c. uptry.

The plate glass is poured melted upon a table covered Chap. II. with a sheet of copper. The plate, as cast, is about an inch thick; but it is ground down to the proper degree of thinness, and then polished. The only manufactory in Britain is at St Helens, about 10 miles from Liverpool. Flint-glass contains much oxide of lead. Dr Lewis extracted from it one-fourth of its weight of that metal in a malleable state . Though it be very solid, it does not seem to be absolutely impervious to gaseous bodies, at least when heated nearly to the melting point. Dr Lewis surrounded a piece of it with charcoal powder, and kept it for some time in a heat not sufficient to melt it. The lead was revived in drops through the whole substance of the glass †. Dr Priestley ascertained, that glass tubes filled with hydrogen gas, and heated, became black, from the revival of the lead. When alkaline hydrosulphurets are kept in glass phials, the inside is soon coated with a black crust. I am informed by Dr Henry of Manchester, that this black crust is nothing else than lead separated by the sulphur from the glass.

Crown-glass is made without lead. It is therefore Crown much lighter than flint-glass. It consists of fixed alkali fused with siliceous sand. As the earthy matters employed by the glass-makers are seldom quite pure from some mixture of iron or similar ingredient, the glass would have a green colour unless some means were taken to remedy it. The addition of some black oxide of manganese remedies this defect; hence it is used for that purpose by the glass-makers, and was formerly

Book II. Division III. called soap of glass. If too much be used, the glass we quires a purple cast; a colour very common in the window-glass made in England. In Scotland the window-glass has always a considerable shade of green.

As no exact analysis has yet been made of the fner kinds of glass, we are not acquainted with the proportion of its constituents. They no doubt vary considerably. As the fixed alkalies are volatilized by a strong heat, it would be worth while to examine whether a portion of them is not driven off while the glass is a fusion, or whether the previous steps of the process prevent that from happening.

Bottle-glass.

Bottle-glass is the coarsest and cheapest kind; little or no fixed aikali enters into its composition. It consists of an alkaline earth, usually lime, combined with alumina and silica. In this country it is composed of sand and the refuse of the soap boiler, which consists of the lime employed in rendering his alkali cause, and of the earthy matters with which that alkali was contaminated. The following are the ingredients used in the bottle-glass manufactory of Lafond in France.

Sand, from......33 to 40 Lixiviated ashes 62 to 55 Sod2...... 5 to 5

100 100

A specimen of this glass analysed by Vauquelin was composed of

Besides a portion of potash so small that if could not be appreciated .

Chap III

Of the different species of glass, the most fusible is flint-glass, and the least fusible bottle-glass. According to the experiments of Saussure; flint-glass melts at the temperature of 19° Wedgewood, crown glass at 30°, and bottle-glass at 47°+; The specific gravity of glass differs considerably according to its constitu-The following are the results obtained by Brisson, as published in the Appendix to Lavoisier's Chemistry:

> Bottle-glass 2.7325 Green glass 2.6423 White glass 2.8922 St Gobin crystal .. 2.4882 -Leith crystal3-1896 Flint-glass 3-9295

Specification vity of glass

The properties that distinguish good glass are well its proper; It is perfectly transparent; its hardness is very considerable; its specific gravity varies from 2.3 to 4, according to the proportion of metallic oxide When cold it is brittle: but at a which it contains. red heat it is one of the most ductile bodies known, and may be drawn out into threads so fine as to be scarcely visible to the naled eye. It is almost perfectly elastic, and of course is one of the most sonorous of bodies. There are but few chemical agents which have any action on it. Fluoric acid dissolves it with great rapidity, and so do the fixed alkalies when as-

^{*} Jour. de Phys. 12. 419.

^{† 1}bid. 2lv. 24:

Book II. Division III. sisted by heat. Dr Priestley has shown also, that the long continued action of hot water is expable of decomposing it: A discovery which explains sufficiently the siliceous earth obtained by Boyle and Margod when they subjected water to tedious distillation in glass vessels.

Method of making glass. After mixing the materials of glass together, it is usual to expose them for some time to a moderate bet. This serves several purposes. It drives off all combustible bodies which may happen to be mixed will the sand; it produces a commencement of combustion which makes the glass afterwards less hable to corrode the clay pots in which it is melted; and it alkali, by this incipient combination, is not so apt to a volatilized; which might be the case if the material were exposed at once to a violent heat. The mixture after being thus heated, is called the frit. Through the domes in which the frit is heated, it is usual to a very thin bubbles of glass passing; a proof that so of the materials are volatilized during this first part the process.

The frit, while still hot, is introduced into large possed of a mixture of pure clay and baked clay, and exposed to a heat sufficient to melt it completely. The fusion must be continued till the effervescence occasioned by the separation of the carbonic acid from the soda has subsided; and the opaque scum, known by the name of glass-gall, which collects on the surface of the glass, must be removed. This scum is occasioned by the common salt and other foreign bodies which are a ways mixed with the soda of commerce. When the fusion has been continued the proper time, the furnity

is allowed to cool a little. In that state the glass is excoedingly duotile, and readily assumes any shape that the workman pleases.

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If the glass vessels, after being formed, were cooled rapidly, they would contract unequally, and become in councequence so brittle as to fall to pieces whenever they were handled. To prevent this inconvenience, they are put into a large red hot furnace, which is allowed to cool very slowly to the temperature of the air. This process is called annealing.

Glass is often tinged of various colours by mixing Coloured with it while in fusion some one or other of the metal-Mc oxides.

Blue glass is formed by means of oxide of cobalt.

Green, by the oxide of iron or of copper.

Violet, by oxide of manganese.

Red, by a mixture of the oxides of copper and iron.

Purple, by the purple oxide of gold.

White, by the oxide of arsenic and of zinc.

Yellow, by the oxide of silver and by combustible bodies.

Opticians, who employ glass for optical instruments, Its defects often complain of the many defects under which it labours. The chief of these are the following:

1. Streaks. These are waved lines, often visible in glass, which interrupt distinct vision. They are probably owing sometimes to want of complete fusion, which prevents the different materials from combining sufficiently; but in some cases also they may be produced by the workmen lifting up, at two different times, the glass which is to go to the formation of one ressel or instrument.

SECONDARY COMPOUNDS.

Book II. Division III.

- 2. Tears. These are white specks or knot, on sioned by the vitrified clay of the farraces, or by the presence of some foreign salt.
- 3. Bubbles. These are air-bubbles which have me been allowed to escape. They indicate want of conplete fusion, either from too little alkali, or the spication of too little heat.
- 4. Cords. These are asperities on the surface of the glass, in consequence of too little heat.

Chap. HL

CHAP. III.

OF SALTS.

THE word SALT was originally confined to muriate of Definition.

and or common salt; a substance which has been known
and in common use from the remotest ages. It was afterwards generalized by chemists, and employed by them
a very extensive and not very definite sense. Every
body which is sapid, easily meited, soluble in water, and
not combustible, has been called a salt.

Salts were considered by the older chemists as a class of bodies intermediate between earths and water. Many disputes arose about what bodies ought to be comprehended under this class, and what ought to be excluded from it. Acids and alkalies were allowed by all to be salts; but the difficulty was to determine concerning earths and metals; for several of the earths possess all the properties which have been ascribed to salts, and the metals are capable of entering into combinations which possess saline properties.

In process of time, however, the term salt was restricted to three classes of bodies; namely, acids, alkalies, and the compounds which acids form with alkalies, earths, and metallic oxides. The first two of these classes were called simple salts; the salts belonging to the third class were called compound or neutral. This Book II. Divates III. last appellation originated from an opinion loog entatained by chemists, that acids and alkaltes, of which they are composed, were of a contrary nature, and the they counteracted one another; so that the resultacompounds possessed neither the properties of acids at of alkaltes, but properties intermediate between the

Chemists have lately restricted the term salt at more, by tacitly excluding acids and alkaltes from a class of salts altogether. At present, then, it denotes only the compounds formed by the combination acids with alkalies, earths, and metallic oxides.

No part of chemistry has been cultivated with me zeal than the salts, especially for these last 40 year. During that time the number of saline bodies has been determined with precision great number have been determined with precision Still, however, this wide and important region is from being completely explored.

Mumber of

Chemists have agreed to denominate the salts from the acids which they contain: The earth, alkalt, a metallic oxide, combined with that acid, is called the date of the salt. Thus common salt being a compound of muriatic acid and soda, is called a muriate, and so is called the base of common salt. Now since there are 34 acids and 65 bases, it would appear, at first sight that there are 2210 salts; but of the 53 metallic or ides at present known there are a considerable number which cannot combine with all the acids. This is

The terms sa't and neutral sait are often confounded. In this West the epithet anarul is confined to salts having no excess of acid or but.

the case also with silica, and perhaps with some of the Chap. III. other earths. We must therefore subtract all these deficiencies from the full number 2210. However, to compensate this, at least in part, there are several acids capable of combining with two bases at once. Thus the tartaric acid combines at once with potash and soda, Such combinations are called triple salts, and they in- Triple salts. crease the number of salts considerably. There are some salts, too, which are capable of combining with an additional dose of their acid, and others which combine with an additional dose of their base. The French chemists denote the first of these combinations by adding to the usual name of the sait the phrase with excess of acid, or by prefixing to it the word acidulous : They denote the second by subjoining the phrase with excess of base. This method has the merit of being precise; bur it is awkward and tedious. The ingenious mode of naming these combinations proposed by Dr Pearson ought certainly to be preferred. It is equally precise, if not more so, and far more convenient in every respect. It consists in prefixing to the usual name of the salt the Supersalts preposition super, to denote an excess of acid, and the sales preposition nub, to denote an excess of base *. Thus sulphate of potash denotes the salt in its state of perfect neutralization, without any excess either of the sulphuric acid or of the potash; supersulphate of potash is the same salt with an excess of acid; subsulphate of potash is the same salt with excess of base. These three different kinds of salts must increase the number of saline compounds very considerably; but the precise number

Postson's Translation of the Chemical Nomenclature, p. 54.

Book II. Division III. of salts is not known, as many of them remain still an examined by chemists. Propably there are not man fewer than 2000. Some idea may be formed of the progress which this branch of chemistry has made, by recollecting that 40 years ago not more than 10 salts a all were known.

Of these 2000, however, a considerable number my be considered as still unknown, as they have been merely formed without being examined. Of those which are known, the greater number have not been applied to any use, and therefore do not deserve a very paramlar description.

Nomenclature. As the different genera of salts are denominated from their acids, it is evident that there must be as may genera as there are acids. The terminations of the names of these genera differ according to the name of the acid which constitutes them. When that acid contains a maximum of oxygen, the termination of the generalise ate; when it does not contain a maximum of oxygen, the termination of the genus is ate. Thus the salts which contain sulphuric acid are called sulphates those which contain sulphurous acid are called sulphates the salts differ very much, according as the acid is saturated with oxygen or not. The iter are seldom permanent; when exposed to the air, they usually attract oxygen, and are converted into ates.

Every particular species of salt is distinguished by subjoining to the generic term the name of its base. Thus the salt composed of sulphuric acid and soda is called sulphate of soda. Triple salts are distinguished by subjoining the names of both the bases connected by hyphens. Thus the compound of tartaric acid, potash,

and sods, is called tartrate of potasb-and-soda. An- Chap. III. other mode of naming these triple salts is sometimes ollowed. One of the names of the bases is prefixed to the generic name so as to act the part of an adjective. For example, soda-muriate of rhodium, means the triple salt composed of muriatic acid, soda, and oxide of thodium. Sometimes the name of the base prefixed is eltered a little, as ammonio-sulphate of magnesia (sulphate of magnessa-and-ammonia); ferruginous sulphate of zinc (rulphate of zinc-and-iron). This is a less unwieldy mode of naming the triple salts, but it is not always possible to employ it in our language.

Before the correction of the chemical nomenclature Arrangeby Morveau in 1781, chemists usually referred the genera of salts to the bases, and distinguished the species by the acids. That celebrated philosopher entirely reversed the method by introducing the new generic terms formed from the acids; and his ingenious nomenclature having been sanctioned and improved in 1787 by Lavoisier, Berthollet, and Fourcroy, who joined with him in forming a new chemical nomenclature, has now become universally prevalent. As far as relates to the salts which have alkaline and earthy bases. this method introduced by Morveau is certainly proper; for in them the acids in a great measure stamp the character of the salt, and therefore ought to constitute the generic distinction. But it does not apply equally well to the salts whose bases are metallic oxides; for in them it is not the acid but the base which stamps the character. In them, therefore, the genera ought to be reversed: they ought to be derived not from the soids but from the bases, or rather from the metals whose oxides constitute the base.

Park R.
Mirana III.
Director de-

The salts, then, naturally divide themsels grand classes; the first of which comprehens line and earthy salts, which derive their means characters from their acids; the second the metalline salts, whose bases, on the countheir most important properties. I shall the wide this Chapter into two Sections: in which I shall describe the alkaline and care the second, the metalline.

SECT. I.

OF ALKALINE AND EARTHY SALT

As the salts belonging to this Section may with great propriety according to their naturally divide themselves into 34 generational acid constituting a genus. But of some must be omitted altogether; because which belong to them have not been examingenus will comprehend under it as many spense bases; besides the triple salts and the appears.

As these genera are very numerous, it tended with considerable advantage to the intuitive them into sets according to their This is attempted in the following TABLE.

TABLE of the Alkaline and Earthy Salts. Incombustible Salts.

Table of the genera.

- a. Not altered when heated with combustibles.
 - 1. Muriates.
 - 2. Fluates.
 - 3. Borates.
 - 4. Phosphates *.
- b. Decomposed without combustion when heated with combustibles.
 - 1. Sulphates.
 - 2. Carbonates.
- c. Set fire to combustibles, or yield oxygen gas by heat.
 - 1. Nitrates.
 - 2. Nitrites.
 - 3. Hyperoxymuriates †.
 - 4. Arseniates.
 - 5. Molybdates.
 - 6. Tungstates.
 - 7. Chromates.
 - 8. Columbates 1.
- l. Combustible Salts.
 - a. Acids partially dissipated, leaving salts in att.
 - 1. Sulphites.
 - 2. Phosphites.
 - b. Acids totally dissipated, leaving the base and charcoal.

Phosphate of ammonia is decomposed by heat and combustibles.

[†] The nitrate and hyperoxymuriate of ammonia are combustible alove. hey disappear completely when heated.

¹ The genera in italics are placed from analogy only.

Book fi. Birisian III.

- 4-Acids partly sublimed unaltered.
 - 1. Acetates.
 - 2. Succinates.
 - 3. Moroxylates.
 - 4. Benzoates.
 - 5. Camphorates.
 - +- Acids wholly decomposed.
 - 6. Oxalates.
 - 7. Mellates.
 - 8. Tartrates.
 - 9. Citrates.
 - 10. Kinetes,
 - 11. Saccolates.
 - 12. Urates.
 - 13. Sebates.
 - 14. Malates.
 - 15. Formiates.
 - 16. Suberates.
- +++Anomalous.
 - 17. Gallates.
 - 18. Prussiates.

Let us take's view of all these genera, following the order of the Table.

As the salts constitute a class of bodies of great importance to the practical chemist, but so numerous that it is difficult to remember the properties of each, it is of some importance to facilitate the means of comparing them together as much as possible. It will contribute considerably to this if the salts belonging to each genus be described exactly in the same order. This shall be attended to as much as possible in the following account of the salts.

The importance of the salts as chemical instruments

is very much connected with their solubility in water. Some are insoluble in that liquid, others dissolve in it with more or less facility. I shall lay hold of this circumstance to subdivide each genus into two parts: the first shall contain the salts which dissolve in water; the second those which are insoluble. Many of the salts have not yet been applied to any useful purpose, while others are of importance either to the chemist or for the purposes of civilized society. After having described the properties of the salts belonging to each genus, I shall add a third part, in which some of the most important uses of the salts shall be mentioned. It will not be expected that I shall indulge in minute details, or attempt to exhaust so fertile a subject. Each genus, then, as far as possible, will be divided into three parts; namely, 1. Soluble salts; 2. Insoluble salts; 3. Uses of the salts of the genus described.

Chap. III.

Book II. Diverson HL

ORDER I.

INCOMBUSTIBLE SALTS.

GENUS I. MURIATES.

The muriates are the salts which have been longest known, and from which indeed the whole of the class have borrowed their name; for to them belongs common salt, the most important and the most indespensibly necessary of all the salts. They may be distinguished by the following properties:

Characters.

- 1. When heated, they melt, and are volatilized, a least in part, without undergoing decomposition. The first portions which fly off contain an excess of acid.
- 2. Not in the least altered by combustibles, even when assisted by heat.
- 3. Soluble in water. For the most part they raise the boiling point of water.
- 4. Effervesce with sulphuric acid, and white acid fumes of muriatic acid are disengaged.
- 5. When mixed with nitric scid, they exhale the o-dour of oxymuristic scid.

The alkaline and earthy muriates at present knows amount to twelve.

L SOLUELE MURIATES,

Sp. 1. Muriate of Potasb.

This salt was formerly called febrifuge or digutive salt of Silvius, and regenerated sea salt.

It may be prepared by dissolving potash in muriatic and, and evaporating the solution till the salt crystales; but it is commonly obtained during other chemiprocesses. Chap. III.

Its crystals are cubes, but often rather irregular.

It has a disagreeable bitter taste. Its specific gravity

1.836 .

Properties.

When boiled in water, it dissolves in 1.7 times its eight of that liquid †. It requires three parts of cold ater 1; but this difference is not sufficient to enable us obtain regular crystals by allowing a saturated boiler solution of this salt to cool. Regular crystals can be obtained by abandoning the solution to sponneous evaporation.

It suffers little alteration from exposure to the air.

When exposed to heat, it decrepitates, melts when exacted to redness, and at last is volatilized in a white moke, but without decomposition. In a red heat it uses 2.86 per cent. of its weight ||.

It is not sensibly soluble in pure alcohol.

Many different analyses of this salt have been published, but they by no means agree with one another. The following Table exhibits the constituents according to the respective analists.

^{*} Kirwan.

⁺ Wenzel's Varwandtiebeft, p. 310.

Bergman, Opurc. i. 134.

[|] Wenzel, Ibid. p. 314.

Book II. Division III. Composi-

		+	‡	5		1
Acid.	21	29	31	34	35	36
Base.	74	63	Ø1	66	65	15/M
Water.	5	8	8			
Total.	100	100	100	100	100	100

These results appear altogether irreconciteable unit each other; but the differences are rather apparent in teal. The quantity of muriatic acid was estimated up precipitating it by means of nitrate of silver, and we say ing the muriate of silver obtained. Now But the formed his estimate by supposing that 100 parts unit ate of silver contain only 11.5 muriatic acid, whereas they seem in reality to contain 18. Had he used the true proportion, the quantity of acid instead of 21 would have been 33. The water was estimated by expension the salt to a red heat; now in this temperature, some of the acid probably escapes as well as the water. Upon the whole, the estimate of Wenzel appears to me to be the nearest to accuracy.

Sp. 2. Muriate of Soda.

History.

This salt has been known and in common use as a seasoner of food from the earliest ages. It is known by

Buchole, Bestrage, isi. 133.

⁺ Wiegleb, Ibd.

Bergman, Opuse, i. 134.

Rose, Gehlen's Jour. vi. jt.

Wenzel, 'erwandinbaft, p. 100. The water is supposed expelli-

Kirwan, Nicholson's Querte Jour. iti. 215.

gem. In this country the term salt is usually aplied to it without any addition. The nature of its acid
as discovered by Glauber, if it was not known earlier.
The affirms in his Specimen Beccherianum, that its base
an alkali; but Duhamel was the first who showhow to obtain that base in a separate state, and who
monstrated the difference between it and potash.

This salt exists in great abundance native, and therehere is never formed artificially by chemists. Immense wasses of it are found in different countries, which rewire only to be dug out and reduced to powder. In hat state it is called rock salt. The water of the ocean Mso contains a great proportion of this salt, to which deed it owes its taste, and the power which it posases of resisting freezing till cooled down to 28.5°. When this water is evaporated sufficiently, the salt prepitates in crystals. It is by this process that it is ob-Mined in this country. But the salt of commerce is not sufficiently pure for the purposes of chemistry, as it conmins usually muriate of magnesia, &c.; but it may be obtained pure either by repeated crystallizations, or by the following process: Dissolve it in four times its weight of pure water, and filter the solution. Drop init a solution, first of muriate of barytes, then of carconate of soda, as long as any ir cipitate continues to Separate the precipitates by finitiation, and evaporate slowly till the salt crystal zes.

Muriate of soda usually crystal...zes in cubes, which, Properties.

^{*} Mineralogy, 1 . 57.

578 SALTS.

Book II. Divn - Iti, of the cubes are truncated; and in urtne the common salt, as Foureroy and Vauquelin I tained, assume the form of octahedrons.

Its taste is universally known, and is willy speaking denominated sale. Its specific 2.125. According to Bergman, it is solutiones its weight of cold water, and in 2. weight of hoiling water †. When it is bottome in water, it dissolves in 2.59 times it that liquid ‡. The specific gravity of the slution is 1.198, temperature 42. Pure not dissolve this salt, but it is somewhat a cohol of .830.

It is not affected by exposure to dry air.

of sods of commerce, indeed, contains a quariate of magnesia, which renders it deligcontains also sulphate of magnesia and sulphate

when heated, it decrepitates. In a red and evaporates in a white smoke without decomposition. In a red heat it loses about of its weight ||. Part of this loss is to be the muriane acid ¶.

The following Table exhibits the result out analyses that have been made to ascerta

BACK & P. THE METHOD

^{*} Fahrenheit, PAR Trees 1784, vol avani 114 Eine 2420, and De Watton found a 2420, See his Faire, v. 6

^{\$ 00}pmin 1 134

Wareto's Glom, Frienz, tal.

⁹ Seignette, Baume's Clemie

		+	1	\$		1	
Acid	28	15 10	40	43:20	-	52	44
Base	55	53.00	58	56.80	54	42	50
Water	6	8-12	2			6	6
Total	100	100	100	100	100	100	100

Chap. 217. Compositien.

small proportion of acid obtained by Bucholz is be ascribed to his mode of estimating the constituas of muriate of silver. When corrected it would beone 44.3, which approaches the result obtained by Rose; and as it is nearly a mean of the analyses of fewan and Bergman, is probably very near the truth. The last column, then, founded on this result, is an apeximation to the constituents of muriate of soda.

Sp. 3. Muriate of Ammonia.

This salt was known to the ancients, and was called History. them sal ammoniac, because it was found in great mantities near the temple of Jupiter Ammon in Afri-10 . Its composition was first pointed out by Geofboy junior in 1716 and 1729 th, and afterwards more recisely by Duhamel in 1735 tt. For many years he whole of the sal ammoniac used in Europe was im-

Bucholz, Beitrage, in. 138.

⁺ Kirwatt, Nicholson's Quarto Jour. id. \$15.

Wiegleb, Bucholz, Ibid. 5 Rose, Gahlen's Jour. vi. 31.

Wenzel, Perwandischoft, p. 101. Bergman, Opnic. i. 134.

De Pliny, lib. axri. c. 7.

¹⁾ Men. Per.



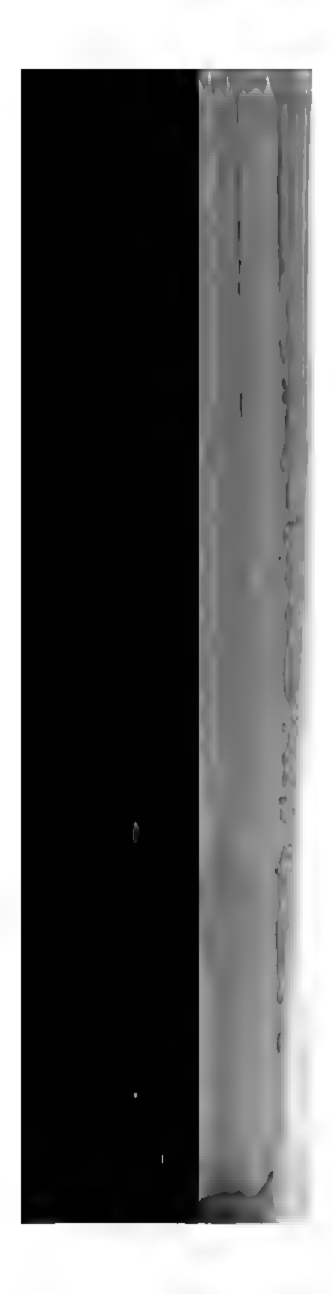
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... or a waste sq

· ----

The state of the s





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this salt is sublimed with gold leaf, there is he neck of the retort an amethyst-coloured ordering on purple, soluble in water, and formple solution. When filtered, there remains purple powder. This salt seems from this to le of oxidizing gold.

Instituents of this salt seem first to have been d by Tournetort in 1700; but he did not sucscertaining their proportion. The result of ent experiments made for this last purpose is 3:

	†	‡	ş	11	1	
i	31	40	42.75	49	76.3	48.
3	69	60	25.00	51	23.7	14.9
er			32.25			37.1
al	100	100	100	100	100	100

Composi-

w estimate of acid in the analysis of Bucholz as has been already observed, upon h statethe proportion of acid in muriate of silver. crected, the true number turns out 48 which

Crell's New Discoveries, &c. Part ii. p. 41.

Beitroge, iii. 146.

Wiegleb, Ibid.

Micholson's Querte Jeur. iii. 216.

Fermendischeft, p. 102.

the result of an analysis, but of a calculation founded in that equal bulks of ammoniscal and muriatic acid gas which is not far from the truth. In the first and amages is included in the proportion of base stated.

mate stated in the sixth column of the preceding like be nearly correct, as is to be presumed, then the war in 100 parts of sal ammoniae must amount to 3 %. From these data, the numbers to the last column of the Table were obtained, which exhibit an approximation to the true proportions of the constituents of this sale Variations in the proportion of the water, which may be expected in different samples, will account for the discordance in the different analyses exhibited above.

Sp. 4. Muriate of Magnesia.

History.

This salt exists in sea-water, and in several mineral springs, particularly that at Lymington in Hampshire. It was first described by Mr Brown in the Philosophical Transactions for 1723 , but its composition was not understood till long after, when the experiments of Black and Margraf had established the peculiar nature of its base. Bergman afterwards published a description of it †.

As it is found native in abundance, it is seldom formed at ificially; but it may be prepared by dissolving carbonate of magnesia in murratic acid, and evaporating the solution to a proper consistency.

Properties.

It is not easily crystallized. Bergman's method was to evaporate it by a considerable heat to the proper degree of concentration, and then to expose it to a sudden cold. By this method he obtained it in small needles 1. Its specific gravity is 1.601 §.

[§] Haucufratz, Aun. de Chim. unvisie 21.



[&]quot; Phil Trine xxxil 348

[;] Bergman, a 382.

[†] Opure. i. 382.

rather more than half its weight of water to dis
t; for when exposed to the air it runs into a liand attracts \(\frac{6.6}{10.6} \) parts of its weight of water.

olubility in alcohol increases with the strength of raid. Alcohol of '817 dissolves half its weight but alcohol of '900 only th of its weight.

if hot water be poured, spongy masses are formeven soluble in muriatic acid 1.

deliquesces very speedily when exposed to the

trong heat decomposes it completely, according accrey. When dried in a high temperature, it is mustic §.

following Table exhibits the proportions of its ments, according to the experiments of different ints:

	11	1	**
Acid	34	34.50	57
Base	41	31.07	43
Water	25	34.34	
Total	100	100	100

Composi-

orgman, Opere. i. 136.

⁺ Kirwan on Mineral Weters, p. 374-

gman, i. 182.

⁽ Westrumb, Ann. de Chem. it 135.

Гезпап, О/ше и 136.

⁴ Kirwan, Nich. Quarte Jour. il. 215.

Wenzel, Vermandtrebaft, p. 104...

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Localing in Fourier. Le manuel é 1º aurose al manuelle 1º aurose al mogretio

W.:

up. s. Marrow of Love.

This said has been long known to chemical It was to homerly called fixed amountar, because it was the north physical by decomposing sai amountaring by the line.

• Opuss. 1. 270. 1 First croy, 111. 370.

† Am. & Clim iv. 222

smally prepared from the residue of that proit may be obtained by dissolving carbonale of muriatic acid.

Chap. III.

systals are six-sided striated prisms, terminated Properties. sharp pyramids. It is not easy to procure it list, in consequence of its extreme tendency to tee. Indeed, in summer, it is next to impo siprocure good crystals; but in winter they form tously, if we dissolve four parts of the salt in water, of the temperature of 70°, and expose the in an atmosphere of 32°.

taste of this salt is very bitter and pungent. Its gravity is 1'76 .

extremely soluble in water; water at 60° diavery nearly four times its weight of it. The ity diminishes very rapidly with the tempera-Hence water at 30° does not dissolve above half untity, while at 1000 it dissolves any quantity hatever. In hot weather, there ore, it cannot be d in crystals, but only in a hard, white, solid mass. Walker has ascertained, that when the evaporathe solution is carried on till its specific gravity 50 at the temperature of 80°, it crystallizes exposed to an atmosphere whose temperature When its specific gravity at 80° is 1.490. mes on cooling the form of a hard pearl-coloured

shol dissolves this salt so copiously when coned, that I presume it is capable of taking up conaly more than its own weight of the salt, even

[&]quot; Hamepfratz, Ann. de Chim. Rvill. 12.

Nicholson's Journal, v. 326.

though previously exposed to the moisture. So much heat lution as to cause the alcohol

When this salt is exposed to ture very speedily, and delique

When exposed to the sotic melts, and then loses its wate violent heat separates a small | thus reduced to a submuriate, ning in the dark, as Homberg has been called the phosphorus

The following Table exhibit ent experiments made to asce this salt, not in the state of cry to a dry mass.

Composi-

	•	
Acid	31	
Base	44	
Water	25	
Total	100	1

These analyses differ very must of Kirwan is nearest the mean, correct.



^{*} Bergman, i. 136. The solt was we to Kirwan. Michelaun's @ to Jun.

[†] Kirwan, Micholash's 2 to Jur. posed to a red heat before ana vasa

Wennel, Verwonstichoji, p. 1 3.

Sp. 7. Muriate of Barytes.

Chap. Hi.

THE properties of this salt were first investigated by Scheele . Dr Crawford wrote a treatise on it in 1790. Since that period many processes have been published for preparing it t. To Kirwan, Hauy, Bucholz, and souillon La Grange, we are indebted for most of the adsitions made to the description of this salt by Scheele.

The easiest method of preparing it would be to dis- Prepara. plve carbonate of barytes in water and crystallize the solution; but as the carbonate of this earth is rare, the alt is usually formed by calcining in a crucible a mixfure of sulphate of barytes and charcoal, decomposing by means of muriatic acid the sulphuret formed, filtrasing the solution, evaporating it till a pellicle begins to form on its surface, then allowing it to cool slowly. The crystals of muriate of barytes gradually deposite. If the salt happens to be contaminated with iron, which is often the case, the crystals are to be calcined, dissolved water, the solution filtered and crystallized again. By this process the oxide of iron is separated. The improvements proposed lately consist in promoting the fusion and decomposition of the sulphate of barytes, by adding to the mixture sulphur, muriate of lime, or common salt.

The primitive form of muriate of barytes is, accord- Properties. ing to Hauy, a four-sided prism, whose bases are equares. Its integrant particles have the same figure 1.

Crell's Assels, ui. 6. English Translation.

See Van Mone, Jour. de Phys. 11v. 297. Goettling, Phil. Mag. xxii.

^{428.} Boutlion La Grange, Ann. de Chew. 21vn. \$39. Resat, Ibid. 1v. 51.

¹ Fourtroy, iti. 232. English Trans.

Book II. Drawill. It crystallizes most commonly in tables. Sometimes a assumes the form of two eight-sided pyramids appeal base to base *.

It has a pungent and very disagreeable taste, miglike all the other preparations of barytes, is possessed. Its specific gravity is 2.8257 to

It requires 2:20 parts of water, of the temperature 190°, to dissolve one of this salt. It is scarcely less to lubic in cold water. Hence it cannot be procured a crystale by cooling a saturated solution of it in hot mater 1.

From the experiments of Bucholz we learn, that puts alcohol while cold dissolves no sensible portion of the but at a boiling heat it is soluble in 400 parts of that he quid \(\). Weak alcohol dissolves a little of it, and the solubility increases with the dilution. Alcohol of 200, for instance, dissolves \(\frac{1}{100} \) of its weight of this silt \(\frac{1}{100} \).

It is not altered by exposure to the open air. When heated, it decrepitates and dries, and when the temperature is very high it melts, but no heat which we can apply is capable of decomposing it.

The following Table exhibits the experiments made by chemists to ascertain the proportion of its constituents.

Crawford.

² Hamenfratz, Ann. de Chim. 22min. 12.

^{\$} Crawford. \$ Beitrager, bit. 34.

[!] Kitwan, on Meneral Waters, p. 274.

	•	+	*	#
Acid	20	22.93	23.8	24
Base	64	62.47	76.2	60
Water	16	14.00		16
Total	100	100.00	100	100

Chap. IIL.'
Compen-

These analyses agree nearly with each other. Mr Kir-wan's, from the great care with which it was made, seems entitled to the most confidence.

Sp. 8. Muriate of Strontian.

THIS salt was first described by Dr Grawford in 1790. It was afterwards examined by Dr Hope, by Klaproth, Pelletier, Fourcroy, and Vauquelin.

It may be prepared by dissolving carbonate of strontian in muriatic acid, or by decomposing sulphuret of strontian by means of that acid. The solution is then to be evaporated to a proper consistence, in order to obtain the muriate in crystals.

Its crystals, according to Hauy, are very long, slen- Properties der, hexagonal prisms . It has a peculiar sharp, penetrating taste. Its specific gravity is 1.4402 .

Three parts of these crystals are soluble in two parts of water at the tempereture of 60°. Boiling water dissolves any quantity of them whatever ¶. From the ex-

[•] Kirwan, Nicholson's Quarto Journal, iii. 25. The salt in the second column was in crystals, in the fourth dried.

[†] Aiken, Nicholson's Jour. xxii. 301.

[†] Fourcroy, iii. 236. Engl. Trans.

[§] Ib. 262.

[#] Hassenfratz, Ann. de Chim. xxviii. 12.

Ascording to Crawford, they dissolve in 0.825 of water of the tem-

periments of Rucholz, we learn that 24 parts of peralection, at the temperature of 60°, dissolve one parts this salt; and that it is soluble in 19 parts of pure buing alcohol. The alcohol solution burns with 1 in purple colour.

The crystal suffers no change when exposed to the air, except it be very moist; in which case they delequesce.

When heated, they first undergo the watery fmin, and are then reduced to a white powder. A very viden heat is said to decompose this sait.

Muriatic acid, precipitates this salt from its solution in water †. Hence it crystallizes most readily when there is an excess of acid in the solution.

Composi-

The constituents of this salt, according to the malyses hitherto given, are as follows:

	‡		1	H _
Acid	18	23-6	31	38.33
Base	40	36-4	69	61.67
Water	42	40		
Total	100	100	100	100.00

Sp. 9. Muriate of Alumina.

History.

This salt was first described by Margraf in his dissertation on the Earth of Alum, published in the Berlin

perature 190°; but they are much less soluble in cold water. Heate they crystallize in cooling.

Buchola' Bitroge, ili. 29.

+ Hope.

! Kirwan, Nicholann . Da rto Jeer. ili. 225.

§ Vauquelin, F verrey, un. 265. Engl. Trans.

Rose, Gehlen's Jur. vi. 33

Memoirs for 1754 . Scarcely any addition has been hade to the facts which he ascertained, except by Buholz + and Wenzel 1.

Chap. III.

It is prepared by dissolving alumina in muriatic acid: It is always in the state of a supermuriate.

It is scarcely possible to obtain it in the form of cry- Properties. sals: it is usually either gelatinous, or in the state of white powder.

It is extremely soluble in water, one part of that liaid being sufficient to reduce four parts of the dry salt the consistence of a syrup . When exposed to the ir, it speedily deliquesces into a liquor of a glutinous posistence.

Pure alcohol, in the mean temperature of the atmophere, dissolves half its weight of this salt; and when piling, it takes up at least two-thirds of its weight, at deposits again a part as the solution cools ¶.

When heated, it melts and loses its acid, while the lumina remains behind in a state of purity | . .

Its constituents, according to the analyses hitherto Composimade, are as follows:

	***	**	††
Acid	19	29.8	27.7
Base	30	30.0	72.3
Water	51	40*2	
Total	100	100	100

⁻ Margraf's Opuec. ii. 118.

[†] Beitrage, ini. 109.

¹ Versuandischaft, p. 111.

[&]amp; Buchols, Ibid. p. 115.

[¶] Marguaf, Ibid.

[[] Ibid. p. 111.

[•] Buchole, Ibid. The second column contains his analysis as he

These analyses are altogether suppose that Wenzel's earth was not dried sufficiently.

Sp. 10. Muria

This salt has been examin quelin +, and Klaproth ‡. It to the nitrate of yttria. Like ficulty, melts in a gentle heat, ry rapidly from the atmosphe lize, but runs into a jelly.

Sp. 11. Muriat

This salt has only been exalittle more—known at presties than that it resembles nitr easily crystallized. When c it forms an agreeable sweet lice

Sp. 12. Muriat

Properties.

This salt was first examinaterwards described more particles formed by none

It is easily formed by pour ly precipitated zirconia. It very astringent; by evaporati parent crystals in needles, whi in the air. Muriate of zircon

20 (1) The second second

serves it; the third the seme analysis,

the witch we not have resulted from a

configuration of acid.

er and in alcohol; to the flame of which it does not Chap. III. rommunicate any particular colour. Heat decomposes t; and it is decomposed likewise by the saliva when Taken into the mouth.

When muriate of zirconia contains a little silica, it forms cubic crystals without consistence, and resembling a jelly. These crystals, when exposed to the air, gradually lose their transparency, and diminish in vosame; and there are formed in the middle of the salt white silky needle-shaped crystals.

Muriate of zirconia is decomposed by sulphuric acid; part of the sulphate precipitates, and part remains dispolved in the muriatic acid. When this soid is driven off by heat, the remainder of the sulphate is gradually deposited: if the evaporation be stopped before the mass be reduced to dryness, it forms a kind of jelly when cold. It is also decomposed by the phosphoric, nitrie, tartarie, oxalie, and saclactic acids, which form with zirconia insoluble compounds that precipitate in white flakes.

The gallic acid poured into muriate of zirconia pro- Action of duces a white precipitate; but a green, bordering on grey, if the zirconia contains iron; and this last precipitate becomes, when dry, of a bright black colour, and resembles China ink. The liquid preserves a greenish colour: new portions of gallic acid produce no farther precipitation; but carbonate of ammonia separates in great abundance a flaky matter of a purplish colour, not unlike that of the leys of wine. From these experiments it follows, that gallic acid has a greater afanity for zirconia than muriatic acid has; and that the gallates of zirconia and iron are soluble in muriatic eid.

galbe acid.

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Book 11. Devision 11 f. Carbonate of potash decomposes muriase of arma, and part of the carbonic acid combines with th

Carbonate of ammonta occasions a precipitate, was is mostly dissolved by adding more carbonate.

Prussiate of mercury produces an abundant prantate, which is soluble in muriatic acid; and whelenacquently is not muriate of mercury.

A plate of zinc, introduced into a solution of maint of zircoma, occasions a slight effervescence; the hour becomes milky, and in a few days assumes the form of a white semitransparent jelly.

Alumina decomposes muriate of zirconia with these sistance of a slight heat; the alumina dissolves, these quor becomes milky, and assumes the form of a jely. When the muriate contains iron, it remains in the solution, and the precipitated zirconia is quite pure. Here then, is a method of freeing zirconia from iron.

THE MURI-

Such are the properties of those muriates that we aluble in water. No alkaline or earthy muriate is a present known that is insoluble in that liquid, of count the second division of this genus does not exist.

THE following Table exhibits the solubility of the muribles in water and in alcohol, and the proportion of the constituents of each, selected from the analysis that seems best entitled to confidence.

^{*} Vauguelin, Aso. de Chro. ERIL 201.

	Solubility Solubility Strength Constituents					
of	Want.	Alcohoi.	Alcohol	Acid.	Base.	Water.
ania	30.11	1.2	*834	100	33	78
esia	151	50.0	*817	100	89.8	99•3
	35.46	0.5	*834	100	114	14
	400	100+x	*800	100	118.3	
Š	33	0-38	*834	100	185-7	
tian	150	4.16	'796	100	216.2	233
es	43	0.00	*796	100	314.5	87
ina	400	50	-796	100	100	135
Name 1						
inz						
nia						

Chap III.
So ubil tv
anil constituents of
murates.

the 12 salts described above, there are only five or hat have been hitherto applied to any useful pur-The following are the most important.

Muriate of potash. This salt is employed by the makers to procure the crystallization of alum, a prepared from the waste leys of the soap-makers but purpose.

Muriate of soda. This is without doubt the valuable of all the salts; the uses to which it is sed are too numerous to be detailed here, and, bethey are very generally known.

III. Usto OPTHE MURIA ATES.

As a seasoner of food it seems to be almost exto health. The inhabitants of maritime district
too easily supplied with it to be sensible of its vabut the case is very different in those countries the
at a distance from the ocean. Wherever it is for
the soil of the inland parts of America, thickewild beasts resort in multitudes. They apply
tongues to the impregnated earth, and gather as a
as will satisfy their wants. These places are a
licks by the Anglo-Americans. So eager are
animals to obtain salt, that they will suffer them
to be caught rather than leave the spot.

It is needless to mention the great consumptions alt in order to preserve animal food from putrelast the application of it to the glazing of pottery, at the preparation of leather; the use of it in metallic in puritying oils, in soap-making, and a hundred

Methods of extricating code from it. Its utility in chemistry is equally extensive. In it alone are muriatic and oxymuriatic acids obtain and from it also, of late, great quantities of sodal been extracted, and introduced with advantage as a stitute for the soda formerly obtained from the contion of vegetables.

The acid is easily extracted from this salt by a of sulphuric acid: But to obtain the alkali at a crate is not so easy. The methods which have his succeeded may be reduced to two.

1. Muriate of soda is decomposed by means of a

* See Phil. Mag. 22. 97.

ecid than soda has. The soda by this process is set at. Chap. III. liberty, and may be obtained by evaporation and crysballization. Barytes and potash would answer this purpose completely; but unluckily these bodies cannot be obtained sufficiently pure, except at an expence which precludes their employment. There are, however, three substances, which are also capable of setting the base of common salt at liberty, and of furnishing soda, either pure or in the state of carbonate. These are litharge, lime, and iron.

When about four parts of litharge and one of com- 1. By limon salt, properly pounded and mixed, are macerated in a little water for several hours, and stirred repeatedby, the muriatic acid gradually combines with the oxde of lead, and forms a muriate, while the soda is left in solution, and may be obtained separately by filtra-Sion and evaporation. The decomposition goes on atill more rapidly if the mixture be heated during the process.

The fact, that the red oxide of lead decomposes muriate of soda, which was first observed by Scheele, has given occasion to much speculation among chemists. Mr Hassenfratz endeavoured to account for it, by supposing that the oxide is combined with carbonic acid, and that therefore it is a case of compound affinity. Mr Curaudau has proved that carbonic acid, instead of promoting, impedes the decomposition; and that, in fact, carbonate of lead is incapable of decomposing muriate of soda. He concludes, therefore, that the phenomenon cannot be accounted for by the commonly received laws of affinity . Vauquelin has proved more lately, that

[·] Ann. de chim. 224.35-



the decomposition by means provided the quantity of that of common salt; that the submuriate of lead totally insc · is not decomposed by alkalie decomposition to the attraction an excess of oxide *. But th pletely for the decomposition, oxide of lead has a weaker aff soda has. Berthollet has at rent anomaly by proving, the mixed with a third, for which divide it between them in pro the quantity of each: And it one of these substances with a aubstance combines with the and takes it completely from insoluble compound being in the mixture, the decomposition again as at first after every p oxide of lead and muriate of the oxide and alkali divide t them, so that some of the co and some muriate of lead for mixture. But this being in altogether insoluble, separate in consequence of which a ne acid between the oxide and t this continues, provided the c ficient, till the common salt i

· Asa. 4 Clis



That the aikali may be extracted from common salt Chap. III. by means of lime, may be considered as a fact for which 2 By lime. likewise we are indebted to Scheele. Cahausen indeed and hinted at it in 1717; but his treatise had been forgotten . Scheele ascertained that a mixture of lime ad common salt, formed into a paste, and placed in a moist cellar, was covered with an efflorescence of soda 15 dayst. In 1782 Morveau and Carny procured a atent from the French government to establish a mainfactory at Croisic for extracting soda from common alt by means of lime. Their process was exactly the ame with that of Scheele, only upon a larger scale. It does not appear, however, that the manufactory was ver established. Berthollet has rendered it probable that the soda which is found abundantly on the west of Egypt, is formed naturally by a similar process 1.

To Scheele likewise we are indebted for the disco- 3. By iron, very that common salt may be decomposed by iron. He observed that a wooden vessel placed in a cellar, and containing brine, had its iron hoops covered with an efflorescence of soda. This induced him to dip a plate of iron into a solution of common salt, and to sussend it in a cellar. After an interval of fourteen days, be found his iron incrusted with soda . The same decomposition takes place also if zinc or copper be subhisted for iron |.

2. The second method of extracting soda from common salt is less direct. It consists in displacing the muristic acid by means of some other acid, which may

^{*} See his Helmont no Extut'.us.

¹ Ann. de Chim, 222111. 3 15.

Athenas, dan de Chim aia. 92.

⁺ Scheele, ii. 15.

Schreie, n 14.

thus the soda is lest behind at last in a state of pure. The acids which have been made choice of are these, phuric and the acetic; the boracic, phosphoric, and arsenic acids might indeed be employed, as they compose common salt in a high temperature. The products in that case would be borate of soda, or the phosphate, or arseniate, of the same base, according to the acid. These salts might be afterwards decomposed by means of time, and the soda obtained separate that these acids are a great deal too high priced to adopt determined their employment.

4. By sulphuric acid and its compounds.

Sulphuric acid may be either employed in a sepiral state, or in combination with bases, when the sain work it then forms can be procured at a sufficiently chem rate. Alum, sulphate of lime, and sulphate of um, have been respectively employed with advantage in decompose common salt, and obtain sulphate of sola Alum is said to have been first employed for that puppose by Constantini, a physician of Melle, near Osmaburg, about 1750. The process, it is affirmed, does not succeed except at a low temperature . Sulphate of lime decomposes common salt when formed with it into balls, and exposed to a strong heat +. Much discussion has taken place among the German chemists about the possibility of decomposing common salt by mem of sulphate of iron. That sulphate of soda may be obtained by exposing a mixture of these two salts to a strong heat, was first announced by Vander Ballen-This was contradicted by Hahneman, but confirmed by

[†] Malherbe and Athenas, Am. & Chin. ziv.



[.] Jour. de Min. No. iii. p. 53.

the experiments of Tul ten *, Lieblein +, and Weglib 1. Chip. III. It succeeded completely with the French commissioners, Lelievre, Pelletier, Larcet, and Gircud, who were appointed in 1794 to examine the different processes for obtaining soda trem common sall. They ascertained also that pyrites or supersulphuret of fron may be employed for the same purpose f.

After obtaining the sulphite of sods, it is necessary to expel the acid in order to procure the soda separately. This is accomplished by calcining the sait mixed with a certain portion of charcoal or of pit-coal. By this process it is converted into sulphuret of soda, and the sulphur may be abstracted by the intervention of iron or chalk. When the sulphuret of soda is nearly in Juston, small bits of iron (the parings of timplate answer best) are thrown in gradually in sufficient quantity to decompose the sulphuret. The fire is raised till the mixture melts. The sulphur, having a stronger affinity for the iron, combines with it and leaves the soda, which may be separated by solution in water, filtration, and evaporation | . Carbonate of lime may be employed also for the same purpose ¶.

Some chemists have proposed to decompose common 5. By accsalt by means of acetate of lead, using either the acetate of commerce, or one formed on purpose, by combining litharge with the acid liquor obtained by distilling wood. The acetate of soda formed by mixing

tate of lead.

[#] Crel's Annals, 1790. ii. 406.

[†] Ibid. p. 509.

^{\$} lbid, 1793, L 204-

[§] Am. de Chim. 212, 58.

Athan, Ann. as Chim. ziz. 77 .- The process was discovered by Malbeibe Sei Juir. de Min. No. 14. p. 67.

¹ Lebianc, Ann. de Chim. BIR. GE.

common sait with these acetates is afterwards casing in order to decompose and expel tile acetic acid. But these salts are too high priced to be employed with advantage to extract soda from common sait.

III. Muriate of Ammonia. This salt is in some de. gree combustible. Hence its importance to preventite oxidizement of metals. For many years the whole d the sal ammoniac used in Europe was imported from Egypt. In that country the greater part of the feel consists of the dung of their cattle formed into balls and dried. These excrementitions matters seem to contain muriate of soda, or rather muriate of ammonia ready formed; owing perhaps to the saline matters on which the animals feed. The soot formed during the combus tion of this fuel is carefully collected and put into large glass bottles, which are exposed, in furnaces constructed on purpose, to a pretty strong heat. The sal ammopiac gradually sublimes, and attaches itself to the up per part of the bottles, where it forms a cake of some inches in diameter. Though this process was commu nicated to the Academy of Sciences in 1719 by Lemere French consul at Cairo, it was a considerable time be fore the chemists in Europe thought of imitating it, or of preparing sal ammoniac themselves. The first manufactory in Germany was begun by Gravenhorst in 1759 †. Soon after it was made in France by Baume, and in Scotland by Dr Hutton.

Prepara-

The processes most commonly followed in Europe

For a detailed account of the processes of making salt followed in different countries, the reader is referred to Browningg's Act of melicine Common Sail, and to Watson's Ghemical Essays, ii. 32.

[#] Wiegleb's Griebiehte, ii. 5 c.

that salt with muriate of soda; and to expose the mixture to a heat sufficient to sublime the muriate of ammonia. The ammonia is usually obtained by the distillation of animal substances, or from soot. 2. To decompose muriate of time by means of ammonia. 3. To combine muriatic acid directly with ammonia, and to sublime.

This salt is applied to a great variety of purposes. It is from it that pure ammonia is usually extracted. A considerable portion of sal ammoniac is consumed by coppersmiths, &cc. who employ it to prevent the oxidizement of the surface of the metals which they are covering with tip. Dissolved in nitric acid, it forms aqua regia, employed in the solution of gold. It has the curious property of rendering many metallic oxides volatile, and is often used by metallurgists to separate metals from each other. These different metallic flowers, as they were called, or combinations of sal ammoniac with metallic oxides, were formerly used in medicine. Great quantities of this salt were once consumed by the dyers, though it is difficult in many cases to see for what purpose. At present, much of the sal ammoniac made in this country is said to be exported to Russia.

IV. Muriate of Lime. This salt was proposed by Fourcroy as a remedy in scrosulous diseases; but it does not seem to have come into use in this country as a medicine. The discovery of Lowitz of the great cold which it generates when mixed with snow, has made it extremely useful as an ingredient in freezing mixtures. The same chemist has taught us the method of purifying alcohol and ether by means of it, from the water and alcohol with which they are respectively contaminated.

Chap, III.

V. Muriate of Barytes. This salt has been tecommended as a cure for scrofulous disorders. The dose is from five to twenty drops or more. Care ought to be taken not to use it in too great quantities, as, like all other barytic salts, it is poisonous.

In chemistry it is much employed as a reactive to detect the presence of sulphuric acid. When dropt into a liquid holding that acid in solution, an insoluble precipitate of sulphate of barytes immediately appears. Bergman informs us that this precipitate is visible when the acid amounts only to 0.0002 of the liquid. Even when only 0.00009 of sulphuric acid is present, a slight cloud appears in a few minutes after dropping in the muriate.

The other muriates have scarcely been brought into common use either in chemistry or the arts.

GENUS II. FLUATES.

THESE salts were first made known to the world by Scheele in 1771 †; and succeeding chemists have done little more than repeat and confirm his experiments.

Characters.

Flustes may be distinguished by the following pro-

- 1. When sulphuric acid is poured upon them, they emit acrid vapours of fluoric acid, which have the property of corroding glass.
 - 2. When heated, several of them phosphoresce.
- s. Not decomposed by heat, nor altered by combus-
 - 4. Combine readily with silica by means of heat.

Bergman, i. 100.

4 Scheele, i. 26.



Most of them are but sparingly soluble in water; but their properties have been but imperfectly examined.

Chap. III.

I. Solvale Fluates.

Sp. 1. Fluate of Potash.

This salt is most readily procured by fusing in a platinum crucible a mixture of fluor spar and carbonate of potash. The mass, digested in water, yields a solution, which, filtered and evaporated, leaves fluate of potash, It has scarcely been examined.

According to Scheele, it does not crystallize, but forms a gelatinous mass almost without taste, which attracts moisture from the air. It dissolves readily in water. When exposed to the fire it melts without any e-bullition *.

Wenzel says, that it forms crystals when free from all admixture of siliceous earth †.

Sp. 1. Fluate of Potash-and-Silica.

When fluoric acid is obtained by the common process in glass vessels, it is always combined with a portion of silica. If into this impure acid a quantity of potash, or the carbonate, sulphate, nitrate, or muriate of that alkali, be dropt, a gelatinous precipitate immediately appears, which, when dried, becomes white like chalk, and separates into small loose grains like the sand of an hour-glass. This powder is a combination of fluoric acid, silica, and potash. Its nature was first ascertained by Scheele.

Scheele on Eluer, i. 26.

[†] Gren's Handbuch, i. 531.

boar C

It has an acid taste, and is soluble in about 150 parts builing water; but is again partly deposited as a minimum cools. When strongly heated, it melts into transparent glass, and loses its acid.

Mr Scheele has shown, that a similar triple salts be formed by using soda, or the salts containing so asstead of potash.

Lime also is capable of combining with these trisalts, and forming with them quadruple salts, composed finoric acid, silica, fixed alkali, and lime †.

Sp. 3. Fluate of Soda.

This salt may be formed as the fluate of potential that the solution is evaporated till a pellicle forms its surface, it yields on cooling small cubic crystal finate of sods. These crystals have a bitter and astigent taste; they do not deliquesce in the sir, and but sparingly soluble in water. Before the blow-puttery decrepitate, and melt into a transparent globule

Sp. 4. Fluate of Ammonia.

This salt may be obtained by applying a sufficient to a mixture of sulphate of ammonia and fluor sp. Fluate of ammonia sublimes; or it may be prepared saturating fluoric acid with ammonia. The solutive yields by evaporation small crystals of fluate of amponia. When heated, it sublimes in the state of a superfinate.

† Fourcroy, iii. 306.

1

Scheele, Creli's Annals, i. 214. Engl. Transl. † Ibid. 219.

Sp. 5. Fluate of Alumina.

Chap. III.

This salt does not crystallize; but assumes, when exporated, the consistence of a jelly. Its taste is astringent, and it contains always an excess of acid.

THE FLUATES of yttria, glucina, and zirconia, have not been examined.

II. Insoloble Fluates-

Sp. 6. Fluate of Magnesia.

This salt may be formed by dissolving carbonate of magnesia in fluoric acid. The salt precipitates in a great measure as the saturation approaches.

It is not soluble in water except there be an excess of acid. In that case, by spontaneous evaporation, it forms hexagonal prisms, terminated by a low pyramid composed of three rhomboidal sides.

These crystals are hardly soluble in water. Alcohol dissolves a small portion of them. Heat does not de-compose them; nor are they decomposed by any acid.

Sp. 7. Fluate of Lime.

This salt exists abundantly native. It is from it indeed that fluoric acid is always extracted.

It is found frequently crystallized. The primitive form of its crystals, according to Hauy, is the octahedron, but it occurs more frequently in cubes; sometimes the angles, and sometimes the edges of these cubes, are truncated. The form of its integrant particles is the regular tetrahedron.

^{*} Bergman, i. 384.

Phosphoresces. This salt has no taste. Its specific gravity is 3'15. It is insoluble in water, and not altered by exposure to the air.

When heated, it decrepitates and phosphoreses strongly in the dark. It emits this light even under water, or in the vacuum of an air pump. When kex hot for some time, it ceases to shine, and the phosphorescent property cannot be again restored to it by my process known, except by decomposing it altogether by means of sulphuric acid, and forming it anew. Scheele ascertained that new-formed fluate of lime is equally phosphorescent with native. The cause of this curious property is not well understood. After being heated the salt, though it refuses to phosphoresce any more, has not lost any perceptible weight, nor is it altered in any of its other qualities. When strongly heated, fluste of lime melts into a transparent glass. According to Saussure, this takes place at the temperature of 510 Wedgewood .

When heated with sulphuric, phosphoric, or arsenic acid, the fluoric acid is driven off in white fumes; but neither the nitric nor muriatic decompose it, according to the experiments of Scheele. The following Table exhibits its constituents according to the most accurate analysis hitherto made.



Chap. III.

	T	‡	
Acid	16	32 ¹	324
Base	57	67;	671
Water	27	0	0
Total	100	100	100

Sp. 8. Fluate of Barytes.

Is salt, as Bergman informs us, may be formed uring fluoric acid into nitrate or muriate of barytes. cof barytes precipitates in the form of a white er, which requires a considerable proportion of to dissolve it.

Sp. 9. Fluate of Alumina-and-Soda.

Is curious compound has been found native in land, and described by mineralogists under the of cryolite. It has the appearance of a stone. colour is greyish white; it has some transparentiated broken its fragments are cubical. It is soften fluor spar, brittle, and of the specific gravity. Before the blow-pipe it melts. According to alyses of Klaproth and Vauquelin it is composed

n's Handbud, i. 533. This analysis is very erroncous. It is as scheele; but I cannot find it in either of the dissertations on cid written by that illustrious chemist. It is given by Karwan, quoting Scheele.

my analysis.

[|] Klaproth, Beitrage, iv. 369.

Acid and water	40	41
Soda	36	32
Alumina	24	21

100 °

100 1

Sp. 10. Fluate of Silica.

in glass vessels, contains always a portion of silica, a forms therefore in reality a superfluxte of that earth. It this solution be allowed to remain for a consideral time in a vessel not completely shut, it deposites substitution to be flusted of silica. These for croy has ascertained to be flusted of silica. The amay be separated by means of heat, and by concern ted acid. This salt is soluble in alkalies, and for with them triple salts? The silical which precipital when fluoric acid prepared in glass vessels is absorbed water, retains some of that acid even after it been dried.

The fluate of strontian has not been examined.

SUCH is an imperfect detail of the properties of a flustes, a set of compounds which have hitherto attraced but little of the attention of chemists. The recodiscovery of fluoric acid, however, as a constituent many bodies in the mineral kingdom, where it was a suspected, will no doubt raise this neglected genus in

Klaproth's Beitroge, in, a La. | Vanquelin, Hany's Mas. ii 400

more repute, and induce some experimenter to favour Chap. III. the chemical world with a more complete detail. Richter indeed seems to have subjected the greater number of them to an analysis. The following Table exhibits the result of his experiments and calculations .

Flustes of	Acids	Best.
	100	
Ammonia	100	157
Lime	160	186
Soda	100	201
Strontian	100	311
Potash	100	976
Barytes	100	520

Constituents of that

III. Usta OF THE FLUATES.

THE only fluste hitherto applied to any use is the fluate of lime. In Derbyshire it is turned into boxes, candlesticks, and various trinkers: it is employed to facilitate the fusion of different kinds of ores. From it all the fluoric acid is procured, to what use soever that acid is to be applied.

GENUS III. BORATES:

THOUGH some of these salts have been long known. and one of them has been in general use for many years, their nature is still but imperfectly understood, because

I have not been able to procure a sight of Richter's very cursous and important writings on Steebessetry, in which his observations on the Austes are to be a card. The table at the feat was calculated from a short table given by Euchlit in Berthollet's Marique Chemique, i. 136.



they have been but superficis is the only chemist, if we exc tempted a description of them guished by the following pro

Characters-

- 1. Before the blow-pipe th
- When their concentrates sulphuric soid, and allowed boracic acid are deposited.
 - 3. They are not altered by
- 4. With most metallic oxi and form globules of coloures

I. SOLVELE BORATES

Sp. 1. Borat

This salt, which was first little known. It may be proture of boracic acid and nitre nitric acid, and leaves a white dissolved in water, yields cry four-sided prisms. It is cap excess of base, and formin From the experiments of We constituents seems to be *.

Acid...... Base.....

Sp. 2. Bora

This salt, which may be f with boracic acid, has never

* Vermandisci



informs us, that about half its weight of boracic acid is Chap. Ht. necessary to saturate borax . Its specific gravity is 1 351 +. It is soluble in 2-5 of water at the temperasure of 147° 1. From the experiments of Wenzel, the proportion of its constituents seems to be \$

Acid......100 Base 44

Sp. 3. Boran or Sub-borate of Soda.

Tires salt, the only one of the borates which has been History. accurately examined, is supposed to have been known to the ancients, and to be the substance denominated corysocolla by Pliny. At any rate, it is mentioned by Geber as early as the ninth century under the name of borax. Its composition was first pointed out by Genffroy in 1732, and Baron in 1748. Bergman was the first who demonstrated that it has an excess of base, and is therefore in the state of a sub-borate.

This salt is brought from the East Indies in an impure state under the name of tinkal, enveloped in a kind of fatty matter, which Vauquelin has ascertained to be a soap with soda for its base. When purified in Europe. it takes the name of borax. The purification is performed by the Dutch; but the process which they follow is not known. Valmont Bomare informs us that they extract 80 parts of pure borax from 100 parts of tinkal. The operations are conducted in leaden vessels. and consist chiefly in repeated solutions, filtrations, and

Bergman, iii. 325. According to Withering, twice its weight is reseasty.-Bergman's Sciegraphia, p. 26. Eug. Trans.

[†] Hamenfrats, Ann. de Chim. Elviii. 12-

[!] Wennel, Ibid. p. 309.

Verwardtichaft, p. 248.

Book II. Division III. crystallizations. Valmont Bomare suspects that the employ lime-water; and Fourcroy has shown that the might be useful in decomposing the soap in which cric borax is enveloped.

Properties.

Borax, thus purified, may be obtained crystallized in hexangular prisms, of which two sides are much breaker than the remainder, and terminated by triangular properties. It is of a white colour. Its specific gravity is 1.740 +. It converts vegetable blues to green. In taste is styptic and alkaline.

It is soluble, according to Wallerius, in 20 times in weight of water of the temperature of 60°, and six times its weight of boiling water.

When exposed to the air it effloresces slowly and slightly.

When heated, it swells, loses about four-tenths of in weight, becomes ropy, and then assumes the form of a light porous, and very friable mass, known by the name of calcined berax; in a strong heat it melts into a transparent glass still soluble in water.

When two pieces of borax are struck together in the dark, a flash of light is emitted 1.

Composition. This salt, according to Bergman, is composed of

39 acid

17 soda

44 water

100 |

[•] Fourcroy, iii. 330.

[†] Kirwan. Wallerius makes it 1.720 (Chemistry, p. 266); Dr Walson, 1.757 (Essays, v. 67).

Accum, Nicholsop's Jour. ii. 28. | Bergman's notes on Scheffer.

Sp. 4. Borate of Ammonia.

Chap. III.

This salt has been described by Wenzel. It forms permanent crystals, which have a considerable resemblance to borax. When heated the ammonia flies off, carrying with it a portion of the acid, but leaving a part of it in a state of purity. From the experiments of Wenzel, its constituents are

100 acid 34 base and water.

Sp. 5. Borate of Strontian,

This salt has only been formed by Dr Hope. It is a white powder, soluble in about 130 parts of boiling water. The solution turns the syrup of violets green †. It is therefore in a state of a sub-borate.

> H. Insolu-Ble Bo-Bates.

Sp. 6. Borate of Magnenia.

BERGHAN formed this salt by dissolving magnesia in boracic acid. The solution proceeded slowly; and on evaporation, the salt was precipitated in small irregular crystals. It is soluble in acetic acid. Alcohol decomposes it. It melts easily in the fire without being decomposed 1. This salt has been found native in the mountain of Kalkberg near Luneburg in Germany. It was first analysed by Westrumb in 1788.

Verwandtschaft, p. 249.

[§] Hope, Edm. Trans. iv. 17.

² Bergman, i. 386.

Book II. DivisionIII. Found mative It is in the form of transparent or opaque whiteey, tals, consisting of cubes, 'aving their edges and four of their angles truncated. It is so hard as to scratch that and to strike fire with steel. Its specific gravity a 2.566. When heated, it becomes electric; and, what is singular, the truncated angles are always positively electric, while the opposite entire ones are negative.

These crystals are insoluble in water, and not should by exposure to the air. When heated, they deceptate: in a red heat they lose their lustre, but do not nensibly diminish in weight. In a white heat they ion out of their weight. When exposed to the action of a very violent heat, they melt into a yellow-coloured glass.

According to Westrumb, they are composed of

73.5 acid

14.6 magnesia

11.9 hme

100.0

Hence the salt was considered as a triple compound of boracic acid, lime, and magnesia; but Vauquelinhat shown, that the lime is foreign, and that the mineral is borate of magnesia †.

Sp. 7. Berate of Lime.

This salt may be formed by mixing together limewater and the aqueous solution of boracic acid, or by boiling together lime and pure borax in water. In el-

[.] Hany, Ann. de Chim. iz. 59, . Nicholson, ii. 120, Second Series.

white powder, tasteless, and difficultly soluble in

Chap, III.

Sp. 8. Borate of Barytes.

An insoluble white powder, which has scarcely been examined, formed by the same process as borate of lime.

Sp. 9. Borate of Alumina.

Tuts salt may be formed by mixing together the solutions of borate of soda and sulphate of alumina. It is said to be scarcely soluble in water, and not to crystallize.

When boracic acid and silica are exposed to a strong heat, they melt together into a transparent glass. This compound has received the name of barate of silica from Foureroy.

The other borates are unknown.

III. Uses OF THE BORATES-

The only salt belonging to this genus that has been applied to any useful purpose, is borax. It is sometimes used in medicine as an astringent. It is used as a flux for metals, and enters into the composition of some of the coloured glass pastes made in imitation of gems; but its great use is to facilitate the soldering of the more precious metals. It is employed also as a flux by mineralogists in essaying the properties of minerals by the blow-pipe.

[·] Bergman, iii. 363.

Book II. Division (III.

GENUS IV. 1

This class of salts was first Margraff. Several of the poexamined by Haupt *, Schlos Westrumb and Scheele; but count of them we are indebquelin. They may be disting properties:

Characters.

- 1. When heated along wi
- 2. Before the blow-pipe globule of glass, which in so others opaque.
- 3. Soluble in nitric scid v precipitated from that solution
- 4. Decomposed, at least pa and their acid, which is sep charcoal and heated to redness
- 5. After being strongly he resce.

The phosphates, like the si with an excess of acid, and fo

The phosphates at present l of which are triple salts. So different states, constituting vs

is in the description of the phosphi works, they have been confounded with guided by the experiments of Vauquels



^{*} De Sale Mirobile Peristo, 1740.

^{\$} Jour. de Madicine, 1776.

I. SOLUBLE PROS-

Sp. 1. Phosphate of Potash.

Or this salt there are two varieties: The first, which is in reality a superphosphate, was first described by Margraff in 1740, and distinguished from the phosphate by Lavoisier in 1774. The second, which is a neutral salt, was also noticed by Margraff in 1745, but its constituents remained unknown till they were lately ascertained by Darracq ‡.

Variety 1. Superphosphate. This salt may be prepared by dropping carbonate of potash into diluted phosphoric acid till all effervescence ceases, and then evaporating the solution. It crystallizes with great difficulty into striated prisms; assuming more readily the form of (a jelly, and when the evaporation is carried farther, becoming dry altogether. Its specific gravity, when dry, is 2.8516 \(\frac{1}{2} \). It is exceedingly soluble in water; and when dry readily attracts moisture from the atmosphere, and is converted into a viscid liquid. When heated, it first undergoes the watery fusion; then allows its water of crystallization to evaporate, and is reduced to dryness. In a high temperature it melts into a transparent glass, which deliquesces again when exposed to the air.

Variety 2. Phosphate. This salt may be formed by mixing together superphosphate of potash and pure pot-

throughout. Indeed there is strong reason to suspect, that the difference hetween these two genera of salts depends more upon other circumstances than upon the nature of the acid.

[·] Opuec. i. 24. + Ibid. p. 162. | Ann. de Chim. al. 270.

[|] Hassenfratz, Ann. de Chim. 22viil. 13-

Book II. Division III. ash, and exposing them to a strong heat in a platinum cracible. A white coloured substance is obtained, which is the phosphate in question. Or we may boil any of the alkaline superphosphates in a solution of possib. A white powder falls, which is the salt wanted. By this last process it was procured by Margraf.

The phosphate of potash is tasteless and insoluble in cold water, but soluble in hot water; and it precipition as the solution cools in a gritty powder. It is extensed by fusible; melting before, the blow-pipe into a transparent bead, which becomes opaque on cooling. It is soluble in natric, muriatic, and phosphoric acids, the solutions are thick, glutinous and adhesive. When sufficiently diluted, the alkalies occasion no precipital in these solutions; but when they are concentrated, a precipitate appears.

According to the analysis of Saussure junior, its com-

acid 35

base 65

100 9

This salt has the property of combining, and of forming a triple insoluble compound with lime.

Sp. 2. Phosphate of Soda.

History.

This salt exists ready formed in urine, and was the first known of all the phosphates. It occupied a good deal of the attention of chemists; and the difficulty of analysing it gave occasion to various hypotheses con-

^{*} Sur la Vegete, p. 311.

cerning its nature. Hellot remarked it in urine; and Chap. Ilt. described it, in 1737, as a salt different from those that had usually been observed. Haupt described it in 1740 under the name of sal mirabile perlatum, or " wonderful perlated salt." It was called perlated from the grey, opaque, pearl-like colour which it assumed when melted by the blow-pipe. Margraf examined it in 1 745, and found it would not yield phosphorus when reated with charcoal as the other salts of urine did, but ascertained that it contained phosphoric seid. Rouelle the Younger analysed it in 1776; and concluded from his experiments that it was a compound of phosphoric acid and soda "; but Mr Proust, being unable to obtain phosphorus from it, concluded that its acid was not the phosphoric, but another analogous to the boraeic +. To this substance, which Mr Proust actually obtained, Bergman gave the name of perlated acid, and Morveau afterwards called it ouretic acid. But Mr. Klaproth soon after analysed it, and proved that

Dr Pearson, who introduced it into medicine as a purgative, gave the following process for preparing it:

it consisted of soda supersaturated with phosphoric

acid 1. Scheele soon after made the same discovery 6.

The acid of Mr Proust, then, is merely phosphate of

soda combined with phosphoric acid, or superphosphate

Dissolve in a long necked matrass 1400 grains of Preparacrystallized carbonate of soda in 2100 grains of water at the temperature of 150°. Add gradually 500 grains

of soda.

Jour. de Med. 1776, Juillet. 1 Crell's damb, 2785, a 238.

⁺ Jaur. de Phys. 1781, i. 145.

⁴ Ibid. ii. 187.

Book IL Division III. the liquor for some minutes; and while it is boiling hot, fittrate it, and pour it into a shallow vessel. In it remain in a cool place, and crystals will continue to form for several days. From the above quantities of materials he has obtained from 1450 to 1550 grass of crystals. Apothecaries usually prepare it from the apperphosphate of lime, obtained from bones by means of sulphuric acid. An excess of carbonate of sods is all ded to separate the lime. The liquid is then filters and evaporated slowly till it crystallizes.

Properties.

eute angles are 60°, and the obtuse angles 120°, to minated by a three-sided pyramid. Its specific grava is 1°333°. Its taste is almost the same with that a common salt. It is soluble at the temperature of 50° in about four parts of water, and in two parts of boding water. This solution crystallizes on cooling; but, i order to obtain the salt properly crystallized, the solution should contain a slight excess of alkali. When exposed to the air, this salt very soon effloresces on the surface. When heated, it undergoes the watery fusion. At a red heat it melts into a white enamel. Before the blow pipe it melts into a transparent globule, which becomes opaque on cooling, and its surface acquires applyhedral figure.

It is not altered by combustibles nor metals. With metallic oxides it enters into fusion, and forms a colour ed globule of glass. Sulphuric, nitric, and muriant acids, decompose it partially, and convert it into more

[·] Hamenfeatz, Aco. de Chim. Raviti. 19.

Chap. III.

shosphate of soda. In this state it is more soluble in water, and not so easily crystallized; but may be obtwined by proper evaporation in the state of thin scales, not unlike boracic acid. It was this superphosphate which Proust obtained, and which he considered as a peculiar acid.

The greater number of earths may be fused along with this salt, and converted into glass.

Sp. 3. Phosphate of Ammonia.

This salt also exists in urine, and seems to have been arst accurately distinguished by Rouelle. It was afterwards examined by Lavoisier in 1774, and still more lately by Vauquelin . It is usually prepared by saturating with ammonia the superphosphate of lime obtained from bones, and evaporating the solution to such a consistency, that when allowed to cool the phosphate of ammonia is obtained in crystals.

It crystallizes in four-sided prisms, terminated by Proposics. equal-sided pyramids. Its taste is cooling, salt, and ammoniacal. Its specific gravity is 1.8051 +. It is soluble in four parts of water at the temperature of 60°, and in rather a smaller proportion of boiling water. It is by spontaneous evaporation that it is obtained in the state of regular crystals. It is not altered by exposure to the air. When heated, it undergoes the watery fusion: it then dries; but if the heat be contianed, it swells up, loses its alkaline base, and the acid melts into a transparent glass. It is the only one of the earthy and alkaline phosphates which can be decom-

⁴ Jour. de l'Ecole Polytechnique.

[†] Hamenfratz, Ann. de Chem. Exviii. 12.

Book II. Division III. posed by hext: hence the reason that it yields planphorus when distilled slong with charcoal.

It is decomposed by the sulphuric, nitric, and meratic acids, and by the fixed alkalies and alkaline carta. It is capable of combining with an additional dos of acid, and of passing into the state of a superphosphase

Sp. 4. Phosphate of Magnesia.

It has lately been examined with much precision by the indefatigable Vanquelin †. It is usually prepared by dissolving carbonate of magnesia in phosphoric and and evaporating the solution gradually till the salt crystallizes; but it may be obtained in large regular crystallizes; but it may be obtained in large regular crystallizes; Mix together equal parts of the aqueous solutions of phosphate of soda and sulphate of magnesia. No apparent change takes place at first; but in a few hours large transparent crystals of phosphate of magnesia make their appearance in the solution.

`

Properties.

Prepara-

Its crystals are six-sided prisms, the sides of which are unequal. It has very little taste; however, it leaves a cooling and sweetish impression upon the tongue. It specific gravity is 1.5489 ‡. It requires about 15 parts of cold water to dissolve it. It is more soluble in boiling water, but it crystallizes in part as the solution cools. When exposed to the air, it loses its water of

crystallization, and falls down in powder. When here

[•] Opusc. i. 390. † Jour. de l'Exole Polysubaique. † Hussenfrats, Ann. de Chim-unvivi. 12.

moderately, it is also reduced to a dry powder. In Chap III. high temperature, it melts into a transparent glass.

Sp. 5. Phosphate of Soda-and-Ammonia.

Though this salt, known to chemists by the names microcosmic salt and fusible salt of urine, was extractad from urine, and examined much sooner than any of the other phosphates, it was long before philosophers were able to form precise notions concerning its nature, be even to obtain it in a state of purity. Margraf was the first who pointed out the method of procuring it pure, and who published a detailed description of its proper-Res. He showed that it contained ammonia, and that it yielded phosphorus; but he did not succeed in discoverbg its whole constituents. Fourcroy was the first who gave a precise account of the proportion of its component parts †. According to him, it is composed 1.....32 acid

24 sods

19 ammonia

25 water

100

The properties of this salt are nearly those of the phosphate of sods and phosphate of ammonia joined together. It answers better than the first of them as a flux; because the beat soon drives off the ammonia, and leaves an excess of acid. Its specific gravity is 1.509 1. When exposed to the air, this salt effloresces, and gradually lo-

[@] Opene. i. 123.

[†] Ann. de Chim, vii. 183.

¹ Hassenfratz, Ann. de Chim. xxviii. 12.

Book II. Division III. ses its ammonia; a fact first observed by the Dute.

Chaulnes. Margraf had observed that the ammonia dissipated when the solution of it in water is evapore.

Sp. 6. Phosphate of Ammonia-and-Magnena.

This salt was first discovered by Fourcroy, found it in a calcalous concretion formed in the of a horse. Since this discovery, Fourcroy and viquelin have observed it also in human urine.

It might be prepared by mixing together solution the phosphates of ammonta and of magnesia in ma the triple sait immediately precipitates in the str a white powder. When urine is allowed to rem considerable time in close vessels, it often deposite salt in regular crystals on the sides and bottom of vessel. These crystals are small four-sided prisms minated by irregular four-sided pyramids. This is tasteless, scarcely soluble in water, and not half be altered by exposure to the air. When heate falls to powder, gives out its ammonia, and in a temperature melts into a transparent globule. distilled along with charcoal, phosphorus is obtain Foureroy has ascertained that the phosphate of an nia-and-magnesia, obtained from the calculous con sions of the horse, is composed of

33 phosphate of ammonia

33 phosphate of magnesia

33 Water

Chap- III. BLE PHOS-PHATES.

Sp. 7. Phosphate of Lime.

OF this salt there are two varieties; the first neutral, e other a supersalt.

Variety 1. Phosphate of Lime. This interesting salt, bich constitutes the basis of bones, was pointed out Scheele and Gahar in 1774: but for the first precise scount of its properties we are indebted to Eckeberg ", carcroy, and Vauquelin +.

As this salt constitutes the basis of bones, it is not preparemessary to prepare it artificially. It may be obtainin a state of purity by the following process: Caltime the bones to whiteness, reduce them to powder, and wash them repeatedly with water, to separate seferal soluble salts which are present. Dissolve the whole in muriatic acid, and precipitate by means of ammonia. The precipitate, when well washed and dried, pure phosphate of lime.

Phosphate of lime, thus prepared, is always in the Properties. tate of a white powder; but it is found native in rerular crystals. In that state it is known by the name of apatite. The primitive form of its crystals is, according to Hany, the regular six-sided prism; and the primitive form of its integrant particles is a three-sid-d prism, whose bases are equilateral triangles: But it vey often assumes other forms. It is destitute of taste, asoluble in water, and not liable to be altered by exposure to the air. It may be exposed to a strong heat without undergoing any change; but in a very violent

Terell's Anneli, 1798, i. 323.

[†] Mein, de l'Instit, il. 274.

Book II. Division III. heat it becomes soft, and is converted into a what, mitransparent enamel, or rather porcelain. According to the experiments of Saussure, a heat of 378° Week wood is necessary to produce this effect. It is said in nitric and muriatic acid without effervesoence, at may be again precipitated from them unaltered by mariate of ammonia.

Sulphuric, nitrie, muriatic, fluoric, and several agetable acids, are capable of decomposing phosphered lime; but the decomposition is only partial. Factory and Vauquelin have ascertained, that there aim are only capable of abstracting 0.40 parts of the law while the remainder continues combined with the photphoric acid, constituting a superphosphate of lime. Heat the reason that phosphoric acid is capable also of decomposing partially the combination of these acids wallime; it abstracts as much of the lime as is sufficient convert it into superphosphate. The constituents to phosphate of lime, according to the most accurate an lyses, are as follows:

Composition.

	†	1	5	11
Acid	30-5	39	41	45
Base	69.5	36	59	55
Water.		25		
Total	100	100	100	100

Jour, de Phys. xlv. 26.

[&]amp; Richter, Statuper Chemipur, i. 136



⁺ Klaproth's Bestrage, ill. 105

Eckeberg, Crell's Annals, 2708, i. 112.

Fourtrey and Vanquelin.

These analyses do not differ much from each other, if Chap. III. we except the first, which doubtless gives the proportion of acid too small ".

Variety 2. Superphosphate of Lime. This variety Preparawas first distinguished as a peculiar compound in 1795 tion. by Fourcroy and Vauquelin. It had indeed been often formed before, but chemists had neglected to examine It is this salt which always remains in the aqueous solution when calcined bones are decomposed by means of sulphuric acid: and it may be formed artificially by dissolving phosphate of lime in phosphoric seid, till the acid refuses to take up any more, and afterwards evaporating the solution till the salt crystallizes.

Its crystals are usually thin brilliant plates, resem- Properties. bling mother-of-pearl, which easily adhere together, and acquire a kind of gluey consistency. Its taste is strongly acid. Water dissolves it, and in a greater proportion when boiling hot than when cold: hence a saturated solution of it in boiling water crystallizes on cooling. It attracts a little moisture when exposed to the air.

When heated, it readily undergoes the watery fusion; then swells up and dries. In a high temperature it melts into a semitransparent glass, which is tasteless and insoluble, and is not altered by exposure to the air. When this salt is heated to redness along with charcoal, its excess of acid is decomposed, and converted into phosphorus, and phosphate of lime remains behind. It

Klaproth indeed elsewhere gives the constituents of this salt exactly as in the last column of the table in the text.



is from this salt that phe but the process of Fourcro posing the superphosphate lead, and afterwards decon by means of charcoal, mu portion of phosphorus.

24

Sp. S. Phosp

THIS salt has hitherto Vauquelin *.

Preparation. It may be prepared eithesid with barytes or carbor together an alkaline phosph barytes. In either case the pitates immediately in the

Properties.

This salt is tasteless, incr in water, and not altered b specific gravity is 1.2667 † melts into a grey-coloured its component parts is unkn



^{*} Jour. de l'Eale 1

[†] Hassenfratz, As

When phosphoric acid is dropt into a solution of Chap. IIL. barytes water, a precipitate of phosphate of barytes immediately falls; but this precipitate is redissolved by adding an excess of acid . Hence it follows, that this salt is capable of combining with an additional dose of acid, and forming a superphosphate of barytes.

Sp. 9. Phosphate of Strontian.

This salt was first formed by Dr Hope; but it was more particularly described by Vauquelin in 1797 †.

Like the former salt it may be formed by dissolving carbonate of strontian in phosphoric acid, or by mixing together nitrate of strontian and phosphate of soda. A white precipitate immediately falls, which is the phosphate of strontian.

This salt is tasteless, insoluble in water, and not al- Properties. terable by exposure to the air. It is soluble in an excess of phosphoric acid; a property which distinguishes it from phosphate of barytes. Before the blow-pipe it fuses into a white enamel, and at the same time emits a phosphoric light. It is completely decomposed by sulphuric acid, but by no other. According to Vauquelin, it is Composicomposed of41.24 acid

58.76 strontian

100.00

Sp. 10. Phosphate of Alumona.

This salt has only been examined by Fourcroy. It may be formed by saturating phosphoric acid with alu-

[·] Fourceoy and Vauquelin, Mem. de l'Inint. il 61.

[†] Jour. de Min. An. vi, p. 13.

Book II. Division PL mina. It is a tasteless powder, insoluble in win Dissolved in phosphoric acid, it yields a gruty plate, and a gummy solution, which by heat is contend in a transparent glass.

Sp. 11. Phorphate of Terra.

This salthas only been formed by Vauquein. When the solution of phosphate of soda is mixed with the maphate, nitrate, or muriate of ytima, phosphate of para precipitates in gelatinous flakes.

Sp. 12. Phosphate of Glucina.

Titls salt has been examined only by Vauquelm. He obtained it by pouring phosphate of soda into the solution of glucina in sulphurie, nitric, or muriane and. The phosphate of glucina is precipitated in the state of white powder. It does not crystailize. It is tasteless, insoluble in water, unless it contains an excess of and, and not liable to be altered by exposure to the sit. When heated strongly, it melts into a transparent glass †.

Properties.

SUCH are the properties of the phosphates as fat as we are acquainted with them at present. Hitherto on

[†] Phosphoric acid and kilica, when mixed together and caposed to a strong heat, melt into a beautiful transparent glass, which is not decomposed either by the action of acids or of alkalics. Fourtrop has given this compound the name of phosphate of silica; but it is essentially different from salts, and ought therefore rather to be ranked in some other class of bodies.



[.] Ann. de Chim. BERVI. 158.

complete analysis of their constituents has been published except by Richter. The following Table exhibits the result of his experiments and observations.

Chap. III.

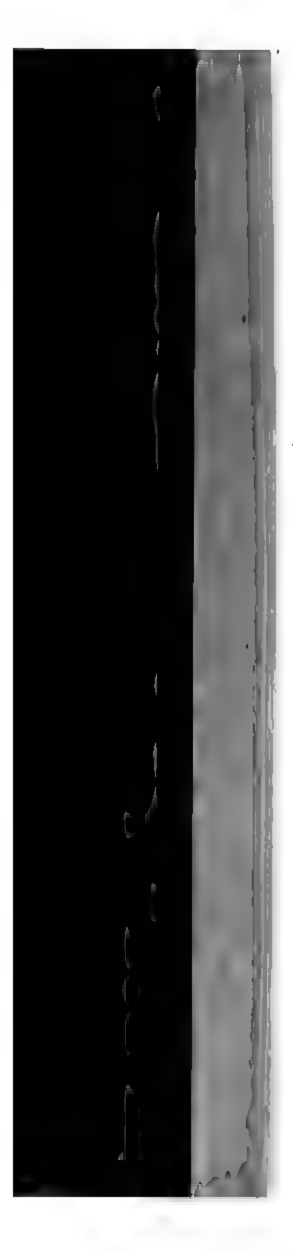
Phosphate of	Acid.	Base.
Alumina	ioo	53.6
Magnesia	100	62.8
Ammonia		
Lime	100	81
Soda	100	87.7
Strontian	100	135.7
Potash	100	164
Barytes	100	227

Four only of the phosphates have been hitherto applied to any useful purpose.

III. Uses of the Phos-Phates.

- 1. Phosphate of soda.—This salt has been introduced into medicine as a purgative; and as its taste is not disagreeable, it has been much employed. It is usually taken in broth, which it is employed to season instead of common salt. It may be substituted for borax to promote the soldering of metals. Mineralogists employ it as a flux when they examine the action of heat on minerals by means of the blow-pipe.
- 2. Phosphate of ammonia.—This salt is much employed as a flux in experiments with the blow-pipe. It enters also as an ingredient in those coloured glasses called pastes, which are made in imitation of precious stones.
 - 3. Phosphate of soda-and-ammonia.—This salt, like

^{*} Statique Chanique, i. 136.



Book 17. Division: Ltf. the preceding, is used in e pipe, and answers remarks readiness with which it par

4. Phosphate of lime. making cupels: from it al phosphorus employed by c employed likewise as a me

GENUS V.

These selts have been I time, and their properties Vanquelin †. They may lowing properties:

1. When heated they en

2. When distilled in a s little phosphorns, and are c

3. Detonate when heate of potash, and are converte

4. Converted into phospl ristic acid.

5. Fusible in a violent !
The phosphites at preser

i. Solvalu Phos-Phites.

Characters.

Sp. 1. Phosp

This salt is formed by di

* The near resemblance between duced me'to place them next each of parate them in the table of the arrai t four, de l' E. de Polytechnique, I.



till it deposites crystals of phosphite of potash. It crystallizes in four-sided rectangular prisms, terminated by dihedral summits. Its taste is sharp and saline. It is soluble in three parts of cold water, and still more soluble in boiling water. It is not altered by exposure to the air. When heated it decrepitates, and then melts into a transparent globule, which becomes opaque on cooling. It does not phosphoresce so evidently as the other phosphites, perhaps because it contains an excess of potash, which saturates the phosphoric acid as it forms.

It is composed of......39.5 acid

49.5 potash

11.0 water

100.0

Sp 2. Phosphite of Soda.

This salt may be prepared exactly in the same way as phosphite of potash. Its crystals are irregular four-sided prisms or elongated rhomboids. Sometimes it assumes the form of square plates, or of plumose crystals. Its taste is cooling and agreeable. It is soluble in two parts of cold water, and scarcely more soluble in boiling water. When exposed to the air it effloresces. Before the blow-pipe it emits a beautiful yellow flame, and melts into a globule, which becomes opaque on cooling.

It is composed of......16.3 acid
23.7 soda
60.0 water

100.0

Snok II.
Division III.

Sp. 3. Phosphite of Aramonia.

This salt may be prepared by the same processes a the two last described phosphites. It crystallizes some. times in long transparent needles, and sometimes in four-sided prisms terminated by four-sided pyramide. It has a very sharp saline taste. It is soluble in two parts of water at the temperature of 60°, and still more soluble in boiling water. When exposed to the air, is attracts moisture, and becomes slightly deliquescent. When distilled in a retort the ammonia is disengaged partly liquid and partly in the state of gas, holding phosphorus in solution, which becomes luminous when mixed with oxygen gas. Before the blow-pipe on charcoal, it boils and loses its water of crystallization; it becomes surrounded with a phosphorescent light, and bubbles of phosphureted hydrogen gas are emitted, which burn in the air with a lively flame, and form a fac coronet of phosphoric acid varour. This gas is emitted also when the salt is heated in a small glass bulb, the tube belonging to which is plunged under mercury.

This salt is composed of 26 acid

51 ammonia

23 water

100

Sp. 3. Phosphite of Ammonia-and-Magnesia.

This salt may be formed by mixing together the aqueous solutions of its two component parts. It is sparingly soluble in water, and may be obtained in crystals; but its properties have not been examined with precision.

Sp. 5. Phosphite of Alumina.

. Chap. IIL

This salt may be prepared by saturating phosphorous acid with alumina, and then evaporating the solution to a proper consistence. It does not crystallize, but forms a glutinous mass, which dries gradually, and does not afterwards attract moisture from the air. Its taste is astringent. It is very soluble in water. When heated, it froths and gives out phosphorus, but it does not readily melt into a globule of glass.

II. INSCEN-BLE PROS-PHITES.

Sp. 6. Phosphite of Magnesia.

This salt is best formed by mixing together aqueous solutions of phosphite of potash or soda and sulphate of magnesia; the phosphite of magnesia gradually precipitates in beautiful white flakes. It has no sensible taste. It is soluble in 400 parts of water at the temperature of 50°, and scarcely more soluble in boiling water. When its solution is evaporated slowly, a transparent pellicle forms on its surface, flakes are deposited, and towards the end of the process small tetrahedral crystals are precipitated. When exposed to the air, it effloresces. When heated, it phosphoresces and melts into a glass which becomes opaque on cooling.

It is composed of.....44 acid

20 magnesia

36 water

100

Sp. 7. Phosphite of Lime.

This salt may be formed by dissolving lime in phosphorous acid. When the saturation is complete, the salt

Book II. Division III. precipitates in the state of a white powder. It is tasted less and insoluble in water; but it dissolves in an excess of acid, and forms a superphosphite. This has salt may be obtained in prismatic crystals by evaporating the solution. This salt is not altered by exposure to the air. When heated it phosphoreses and emits a little phosphorus. In a violent heat it melts into a transparent globule.

It is composed of 34 acid

51 lime

15 water

100

Sp. 8. Phosphite of Barytes.

This salt may be formed by pouring phosphorons acid into barytes water, or this last water into a solution of phosphite of soda. In either case phosphite of barytes precipitates in the form of a white powder. It is tasteless, and but very sparingly soluble in water unless there be an excess of acid. It is not altered by exposure to the air. Before the blow-pipe it meits and is surrounded with a light so brilliant that the eye can scarcely bear it. The globule which it forms becomes opaque as it cools.

It is composed of 41'7 acid

51.3 barytes

7°0 water

100.0

Hitherto none of these salts have been applied to any useful purpose.

GENUS VI. CARBONATES.

Chap. III.

Though several of the carbonates were in the hands of chemists, and employed by them in the greater number of their experiments, their nature and composition were entirely unknown till Dr Black discovered it in 1756. Since that time they have been examined with great attention by almost every chemist of eminence; to that at present no family of salts is more accurately understood. The first treatise on the subject was published by Bergman in 1774.

They may be distinguished by the following proper-

1. When sulphuric acid is poured upon them, they Characters offervesce violently, emitting carbonic acid gas.

- 2. When heated strongly, the carbonic acid is driven off, and the base remains in a state of purity. Some carbonates require a very violent heat to be thus decomposed; but the operation is facilitated by mixing them with charcoal, which decomposes the carbonic acid altogether.
- 3. The alkaline carbonates tinge vegetable blues green, and have an alkaline taste.
- 4. The alkaline carbonates are soluble in water; the carbonates with bases of the alkaline earths are insoluble, but dissolve when an excess of acid is added.

Many of these salts exist native; but they may be all formed artificially by dissolving or diffusing the base in water, and making carbonic acid gas pass into the liquid till it be saturated. At present 11 species are

[@] Opurc. i. L.



known, several of which :

L SOLUBLE CARSO-RATEL

Sp. 1. Gar!

This salt has been long fore its composition was characterized by a great v the manner of preparing i tartar, vegetable alkali, a described with precision by are two varieties of it; th taining an excess of alkali

Preparation. Variety 1. Carbonate.
rating potash with carbonic exposing a solution of potash carbonic acid gas, or by carbonic acid ga

Ртиренцев.

It crystallizes, according lar prisms; the apexes of inverted triangles, converg According to Pelletier the

[†] Ber



^{*} Opus i, 13.

bas eight faces, two hexagons, two rectangles, and four rhombs. It has an alkaline, but not a caustic taste, and still gives a green colour to vegetable blues. Its specific gravity is 2.012+. It is soluble at the common temperature in about four times its weight of water ‡. Boiling water dissolves the of its weight §. Alcohol, even when hot, does not dissolve above trooth part of it. Pelletier has observed, that when the crystallized salt is dissolved in boiling water, bubbles of carbonic acid gas are emitted. It is not altered by exposure to the air. Heat deprives it of its water and part of its acid, but does not decompose it completely.

The constituents of this salt, according to the most accurate analyses hitherto made, are as follows:

		1		++ :
Acid	20	2015	43	43
Base	48	73.5	40	41
Water	32		17	16
Total	100	100	100	100

The coincidence between the experiments of Kirwan and Pelletier is very great, and induces us to confide

[·] Ann. de Chim, xv. 29. † Hamenfratz, Ann. de Chem. zxvul. 12.

[†] Bergman, i. 13. | Pelletier. | Bergman, Open. i. 14.

Richter, Statique Chimeque, i. 136.

^{**} Kirwan, Nicholson's Quarte Jour. in. 215.

[†] Pelletier, Ann. de Chim wv. 33.

Bork II. DivinionIII. much more in their results than those of the other demists.

Prepara-

Variety 2. Subcarbonate. The potash of comme always occurs in this state; but its purity is destroy by the mixture of foreign substances. Pure subezel nate of potash may be procured by heating neutral bonate, prepared by the process above described, to a ness, in a silver or platinum vessel. A portion of 👛 carbonic acid is driven off. The subcarbonate formed has a stronger alkaline taste, and acts with mienergy on animal and vegetable substances that he carbonate. When exposed to the air it soon deliques and assumes the consistency of an oil. It does not in sorb carbonic acid sensibly from the atmosphere, for excess of alkali cannot be removed by treating the with alcohol. Dr Wollaston has shown that this contains exactly one half of the acid which cause carbonate of potash ...

Composition of potush of commerce.

The potash of commerce is always in the state of subsalt; but it contains likewise several foreign a stances which render the proportion of alkali variable. Mr Kirwan has pointed out a very ingenious method detecting the quantity of alkali in any specimen, by a property which it has of precipitating alumina for alum; and Vauquelin has published a still simpler method, namely, the quantity of nitric acid of a given density necessary to saturate a given weight of the sale From his experiments, we learn that the following kind of this salt known in commerce contain the following redients.

[•] Wallastan, On apperaised a † Ann, de Chim, xl. 273.

	Pocash,	Suiphate of potash.	Munare of potash.	Insoluble re-	Carbenje and sod water	Tutal.
Potash of Russia	772	65	5	56	254	1152
America	857	154	20	2	119	1152
American Pearl-ash	754	80	4	б	308	1152
Potash of Treves	720	165	44	24	199	1152
Dantzie	603	152	14	79	304	1152
Vosges	444	148	510	34	304	1440

Sp. 2. Carbonate of Soda.

This salt has been also very long known. It is usually obtained by burning and lixiviating marine plants, or by decomposing common salt. In commerce it is called barilla or soda. In that state, however, it is never perfectly pute, containing always a mixture of earthy bodies, and usually common salt; but it may be purified by dissolving it in a small portion of water, filtrating the solution, and evaporating it at a low heat, skimming off the crystals of common salt as they form on its surface. Of this, like the preceding, there are two varieties; the first neutral, the second containing on excess of alkali.

See a detail of the different plants from which this substance is proved, and of the method of procuring them, in the Amales de Chimie, who sets.

Book I. Divosion III. History. Variety 1. Carbonate. Though this salt occurred tive in Africa in considerable quantities, it will not a tinguished from the soda of commerce, which make carbonate, possessing very different properties, the 1802, Klaproth published an account of it in the variety volume of his works. It is found in the province to Sukena near Fezzan, and is called *trona* by the name. It is crystallized in hard streated masses, not altered by exposure to the air, and of such hardness that the walk of Casse, a fort now in ruins, are said to have been built of it.

This carbonate may be formed artificially by repsing the common subcarbonate of soda to an atmospher of carbonic acid gas. In that situation it does not desssite crystals like subcarbonate of potash, but forms tolid shapeless mass, having some resemblance to the African carbonate.

When the carbonate of soda is exposed to a red bea, it loses its water of crystallization, and likewise a partion of its acid. Its constituents, as determined by the experiments of Klaproth, are as follows:

Acid 39

Base 38

Water 23

100

Variety 2. Subcarbonate. What is usually called the carbonate of soda is increally this salt. It crystalizes in decahedrens, composed of two four-sided pyram ds applied base to base, and having their apexes truncated f.

· Robber . 3.

† Bergman, i. 19.



Chap. HI

It is often obtained also in large transparent flat rhomsoidal prisms. Its taste is precisely the same with that of the carbonate of potash. Its specific gravity is 1.3591. It is soluble in two parts of cold water, and in rather less than its weight of boiling water. So that when dissolved in botting water it crystallizes as the solution pols. When exposed to the air, it very soon efflobesces and falls to powder. When heated, it underpoes the watery fusion; indeed the soda of commerce pometimes contains so much water of crystallization, that, when once melted, it remains permanently liquid. If the heat be continued, the water gradually evaporates, and the salt becomes dry. In a red heat it melts into transparent liquid. A very violent heat drives off . part of its acid. This salt melts rather more easily than carbonate of potash, and for that reason it is preferred by glass manufacturers.

Its constituents, according to the analyses of it hitherto made, are as follows:

	+ 1	1	5	Ś
Acid	16	16	14.42	40-14
Base	20	22	21.58	59*86
Water	64	62	64	
Total	100	100	100	100

^{. ·} Hassenfratz, Ann. de Chim. xavin 22.

[†] Bergman, Opnie. i. 18; and Fourcroy, Systems de Connoissantes, iv. 56.

² Klayroth, in. 65 Engl. Trans.

[&]amp; Kiewan, Nicholeon's Jour. in. 225.

Book II. Divin mill But either these analyses or that of the carbonal a maccurate, or the salts examined were impute. It Dr. Wollaston has shown, that when a pure carbonal of soda is exposed to a red heat it is converted as a subcarbonate, and loses exactly one half of its acid.

Sp. 3. Carbonate of Ammonia.

Prepara-

This salt has been also long known. It is often abtained by distilling animal substances; but for them, cal purposes it is best to extract it from sal ammoniae by means of chalk. Two parts of chalk and one part of sal ammoniae, both as dry as possible, are mixed ingether and put into an earthen retort. On the application of a sufficient heat, carbonate of ammonia substance, and is obtained in the state of a white crystalance mass. This salt, like the other alkaline carbonates, is of two kinds, the neutral and subcarbonate.

Properties.

Variety 1. Garbonate. This salt may be obtained by exposing the con mon subcarbonate of ammonia to the air, or by causing a current of carbonic acid gas to pass through a solution of it in water. It crystallizes to six-sided prisms, usually small, has no smell, and less taste than the subcarbonate. According to the experiment of Schrader, it is composed of about

56 acid

19 base

25 water

100 t.

Variety 2. Subcarbonate. This salt may also be ob-

Phil. Trees, 1808.

Gehlen's Jeer, is

rained in crystals, but they are so small and so irregular, that it is difficult to ascertain their form. According to Bergman, they may be obtained in octahedrons, having, for the most part, their two opposite apexes truncated . The taste and smell of this salt, though much weaker, are the same with those of pure ammonia. Like all the alkaline carbonates, it converts vegetable blues to green, precisely as pure alkalies do. Its specific gravity is 0.966 +. It is soluble in rather less than twice its weight of cold water. Hot water dissolves its own weight of it. Boiling water cannot be employed, because at that heat the carbonate is volatilized. It is not altered by exposure to the air. When heated it evaporates very specdily.

Mr Davy has shown that the component parts of this Companisalt vary exceedingly, according to the manner of preparing it. The lower the temperature at which it is formed the greater is the proportion of acid and water which it contains; and, on the other hand, the higher the temperature the greater is the proportion of alkali. Thus carbonate of ammonia, formed at the temperature of 300°, contained more than 50 per cent. of alkali, while carbonate formed at 60°, contained only 20 per cent. 1. Its constituents of course must be variously stated by chemists, according to the state in which they The following table exhibits some of these found it. results.

[·] Bergman, i. 21. + Hassonfratz, Ann. de Chim. xxvini. 12

Davy & Researches, p. 75.

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	‡	•	H	4
Acid	45	48 to 50	50 to 52	68.2
Base	43			31.5
Water	12		•	
Total	100			100

Sp. 4. Carbonate of Ammonia-and-Magnesia.

This salt has been formed by Fourcroy by mixing together aqueous solutions of its two component parts; but it properties have not been examined.

II. INSOLU-BLE CAR-BONATES.

Sp. 5. Carbonate of Magnesia.

This salt has been long known. Its composition was first discovered by Dr Black; afterwards its properties were investigated by Bergman and Butini, and more lately by Fourcroy*. There are two varieties of it: the first neutral; the second containing an excess of base.

Prepara-

Variety 1. Subcarbonate. This salt is usually prepared by mixing together the solutions of sulphate of magnesia and carbonate of potash, and applying heat. The carbonate of magnesia precipitates in the state of a white powder. Such is the state in which it occurs in

[!] Bergman, i. 21.

[&]amp; Schrader, Gehlen's Jour. ii. 583. When in a solid mass.

[¿] Ditto. When crystallized.

Mirwan, Nicholson's Quirto Jour. iii. 215.

^{*} Inn. de Chim. ii. 278.

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commerce; but Fourcroy has shown that in that state if does not contain a maximum of acid. It is therefore only a subcarbonate; but it may e saturated with acid by diffusing it in water, and making carbonic acid gas pass through it till it be saturated and dissolved.

Variety 2. Garbonate. The carbonate, thus formed, yields by evaporation crystals which are transparent hexagonal prisms, terminated by a hexagonal plane: these are partly in groups and partly solitary: their length is about six lines, their breadth two *. may be obtained in more regular crystals, by mixing together 125 parts of sulphate of magnesia and 136 parts of carbonate of soda, both dissolved in water, filtering the solution, and then setting it aside. In two or three days the carbonate of magnesia crystallizes.

This salt has little taste. Its specific gravity, when Properties. in powder, is only 0.2941 according to Hassenfratz +. It is soluble, when crystallized, in 48 parts of cold water; but when in powder, it requires at least ten times S much; and what is very remarkable, it is more soluble in cold than in hot water impregnated with carbonic acid ‡. When exposed to the air, it effloresces, and falls into powder §. When heated, it decrepitates, falls into powder, and is decomposed.

The constituents of this salt are as follows:

^{*} Butini, sur le Magnesie.

[†] Ann. de Chim. xxviii. 12.

[.] Butini.

Fourcroy, Ann. de Chim. ii. 298.

B: k ii. Divina iil.

	•	+	1
Acid	50	30	36
Base	25	45	43
Water	25	25	21
Total	100	100	100

But the carbonate examined by the two last chemiss does not seem to have been fully saturated with aid. The magnesia of commerce, according to Kirwan, is composed of.........34 acid

45 magnesia 21 water

100

Sp. 6. Carbonate of lime.

This substance, under the names of marble, chalk limestone, &c. exists in great abundance in nature, variously mixed with other bodies. It is perhaps the most important and most generally used of all the salts, unless we except muriate of soda. Its properties of course have been very completely investigated.

Properties.

It is often found crystallized and perfectly transparent. The primitive form of its crystals is the rhomboidal prism with angles of 101^{+0}_{1} and 78^{+0}_{2} . Its integrant particles have the same form. But, besides the primitive, no less than 616 varieties of its crystals have been dis-

Fourcroy and Kirwan.

[†] Bergman.

¹ Burier.

[&]amp; Nicholson's Jour. iii. 215.

any taste. Its specific gravity is about 2.7. It is insoluble in pure water; but water saturated with carbonic acid dissolves \(\frac{1}{1500}\)th part of it; from this solution it gradually precipitates, as the acid leaves it, in the form of a white powder \(\Phi\). It suffers little or no alteration by being exposed to the air. When exposed to heat, it decrepitates and loses its water, and afterwards its acid separates as the heat is increased: But to separate the acid completely, a pretty strong heat is required.

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Its component parts are as follows:

	†	‡	ş
Acid	34	45	50
Base	55	55	50
Water	11		
Total	100	100	100

Composi-

Some very interesting experiments on the suibility of this salt were made by Sir James Hall. The result was, that when the carbonic acid is prevented from making its escape by strong compression, the salt melts at a red heat, and assumes an appearance which has some resemblance to granular limestone. A portion of the carbonic acid is usually dissipated. This portion is

[#] Bergman, i. 26.

[†] Bergman, Opusc. i. 23.

[‡] Kirwan, Nicholson's Journal, iii. 215.

In this analysis the water is confounded with the acid. This wate seems in crystallized pure carbonate to amount to three or four per cent : but I have not been able to ascertain its proportion, because the whole of it separates in combination with the carbonic acid.

' Book II. Division III.

sometimes very small, and very often it does not co. ceed 4 or 5 per cent. Bucholz has lately verified these experiments in a very unexpected manner. Be put 41 pounds of washed chalk (carbonate of lime with only 0.005 of foreign matter) into a crucible, presed it strongly down, and exposed it covered to a strong heat in a furnace. The chalk, except a small portion on the surface, was converted into a foliated, hard. yellowish mass, having considerable transparency, which obviously had undergone a commencement of It was obviously in a similar state with Sir James Hall's carbonate of lime, and contained 42 per cent. of carbonic acid*. Here the same effect was produced without compression. It must have depended no doubt upon the degree of heat to which the crucible was exposed.

Sp. 7. Carbonate of Barytes.

This salt seems first to have been examined by Bergman. Withering first found it native in 1783. Hence it has received the name of Witherite. Since that time it has been examined by Kirwan, Klaproth, Hope, Pelletier, Fourcroy, and Vauquelin, and its properties described with great precision. It may be prepared artificially by exposing barytes water to the open air, or by passing carbonic acid gas into it. In either case the carbonate precipitates in the state of a white powder.

Properties.

It is found native crystallized. Its crystals have been observed to assume four different forms; double

[#] Gehlen's Jur. 2d Series, i. 271.

 $\dot{\Xi}$

six-sided and double four-sided pyramids, six-sided columns terminated by a pyramid with the same number of faces, and small radiated crystals half an inch in length and very thin, appearing to be hexagonal prisms rounded towards the point. It has no sensible taste; yet it is poisonous. Its specific gravity, when native, is 4.331; when prepared artificially, it scarcely exceeds 3.763. Chap. III.

Cold water dissolves $\frac{1}{4304}$ part, and boiling water $\frac{1}{4304}$ part of this salt. Water saturated with carbonic acid dissolves $\frac{1}{430}$ part ¶. It is not altered by exposure to the air. It is decomposed by the application of a very violent heat ||, either in a black lead crucible, or when formed into a paste with charcoal powder.

The constituents of this salt, according to the experiments of different chemists, are as follows:

	•	†	**	‡	9
Acid	7	20	21.67	22	22
Base	65	80	78.33	62	. 78
Water	28			16	
Total	100		100.00	100	100

Compesition.

Sp. 8. Carbonate of Strontian.

This salt was first pointed out as distinct from the History. last species by Crawford in 1790; but we are indebted

[¶] Fourcroy, Ann. de Chim. iv. 64.

Dr Hope.

^{*} Bergman, Opusc. i. 22.

[†] Withering.

^{*} Aiken, Nicholson's Jour. xxii. 303.

¹ Pelletier.

[§] Kirwan, Nicholson's Quarte Jour. iii. 215.

Book II. Divinon III. for the first accurate account of it to Dr Hope. It experiments were afterwards confirmed by Klapes Pelletier, Fourcroy, and Vauquelin. It has been for native at Strontian in Argyleshire and at Leadhilla Scotland. It is usually in streated semitranspan masses, which have a greenish tinge.

Properties.

It has no taste. It requires 1536 parts of boild water to dissolve it *. Its specific gravity is about to It is not altered by exposure to the air. When we ly heated in a crucible, it loses part of its and, to this decomposition is facilitated by making it into paste with charcoal powder. According to Saussi it melts into a transparent glass at the temperature 226° Wedgewood †. When thrown in powder to burning coals, it produces red sparks.

Its component parts are as follows:

Company

	1	5	11
Aeid	30-2	30	30.0
Base	61.2	62	0913
Water	8.0	9	0.2
Total	100.0	100	10(*0

Sp. 9. Carbonate of Alumina.

THE greater number of chemists have agreed in a mitting the existence of this salt. Bergman could not

[.] Hope, Trans. Edin iv 5.

⁺ Jour. de Phys 21v. 24.

Hope, Teans. Edin. iv 8.

S Pelietier, Ann. de Glom. BRI 136

[|] Klaptoth, Beitrage, i. 270. and Kiewan, Nicholson's Jour. up 215

Chap. HI.

when alum is mixed with an alkaline carbonate, part of the alumina remains in solution till the carbonic acid be driven off*. Saussure has shown lately, that water saturated with carbonic acid, is capable of dissolving alumina; but this combination is destroyed by simple exposure to the air. Carbonate of alumina, then, cannot exist in a dry state. What had been considered formerly as a dry carbonate is a triple compound of alumina, carbonic acid, and the alkali employed in precipitating the alumina †.

Sp. 10. Carbonate of Yttria.

This salt may be formed by precipitating yttria from its solution in acids by means of an alkaline carbonate. It is a white, tasteless, insoluble powder, composed of

18 acid

55 yttria

27 water

1001

Sp. 11. Carbonate of Glucina.

This salt has been examined only by Vauquelin. It may be prepared by precipitating glucina from its solution in acids by an alkaline carbonate, and washing the precipitate sufficiently with pure water. It is in the state of a white soft powder, which has a greasy feel. It has no taste, and is exceedingly light. It is insolu-

Bergman, i. 32.

[†] Jour. de Phys. lii. 28.

^{\$} Klaproth's Beitrage, iii. 67.

Book II. Division III. ble in water, not altered by exposure to the air, easy decomposed, and its acid driven off by the application of heat.

Sp. 12. Carbonate of Zirconia.

This salt was formed by Vauquelin by precipitating zirconia from its solution in acids by alkaline carbonates. It is a tasteles white powder, composed of

44.5 acid and water 55.5 zirconia

100.0

When heated the carbonic acid is driven off. This salt is soluble in the three alkaline carbonates, and seems to form with them triple salts.

Such are the properties of the carbonates. The following Table gives a view of their constituents according to the most accurate analysis:

Carbonates of	Solubility	-	Constitue	nts.
Carbonateror	Water	Acid.	Base.	Water.
Ammonia	50	100	33.9	44.0
Magnesia	2	100	50	50
Potash	25	100	95.8	37
Soda	50	100	97.4	59
Lime	0	100	122	
Strontian	0	100	231	
Yttria	0	100	305'5	150
Barytes	ρ	100	354.5	

Table of the constituents of the carbon hates.

All the carbonates are insoluble in alcohol; but the alkaline subcarbonates are partially soluble in that liquid:

III. Uses OF THE CARBO-

From the peculiar nature of the acid which exists in these salts, the alkaline carbonates were long confounded with their bases. The alkaline character of these bases is by no means destroyed by the presence of the acid, and it is easily removed either by means of heat, or by the action of some other base. Hence the alkaline carbonates are applied to all the uses for which the pure alkalies are adapted. These are too numerous and well known to require a particular detail. The use of carbonate of lime as a manure, a paint, and a medicine, ander the names of lime, whiting, crab's eyes, &cc. is

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Uu

Book II. Division III. have come into use, except the mix as a sia, which is occasionally employed as a

GENUS VII. SULPRATEL

Surphyric Acto is capable of capable at the alkalies and earths except silics. The almost all capable of assuming a crime. Their taste, when they have any, is almost a the following properties.

Maracters.

1. They are insoluble in alcohol. Was dissolved in water, alcohol precipitates them crystallized.

this there are numerous exceptions. Most of a sulphates are decomposed by a red heat enter or totally; part of the acid escapes unaltered as has been shown by the experiments of Gay. It is decomposed into sulpharous acid and oxygen parated in the same way, partly unaltered, parated in the same way, partly unaltered, partly unaltered, parated of sulpharous acid and oxygen gas.

3. When heated to redness along with charea

4. When barytic water, or a solution of any a taining barytes, is dropt into a solution of ary sulphates in water, a copious white precipitate diately falls, which is insoluble in acetic seid.

* Men. PAprell, Larg.

he genus of sulphates contains more species than other, owing to the great tendency which the acid to form triple salts. Many of the sulphates are casle of combining with an excess of acid, and of ming supersalts.

Chap. III.

I, Solvale SUL-PRATES.

Sp. 1. Sulphate of Potasb.

OF this salt there are two varieties. The first is natral, the second contains an excess of acid.

Variety 1. Sulphate. This salt may be formed by curating diluted potash with sulphuric acid, and then aporating the solution gently till crystals appear. It ems to have been known at a very early period by semists, and a great variety of names were given to it, scording to the manner of forming it, or the fancy of be operator. Some of these names were, specificum Names. wgans, nitrum fixum, arcanum duplicatum, panacea boltica, sal de duobus, sal polychrest glaseri, &c. but it as commonly known by the name of vitriolated tarr till the French chemists called it sulphate of potasb hen they formed their new nomenclature in 1787.

When the solution of sulphate of potash is sufficiently Properties. duted, it affords by evaporation hexahedral pyramids, short hexangular prisms, terminated by one or more exangular pyramids. But these crystals vary much in heir figure, according to the care with which they are repared.

It has a very disagreeable bitter taste. Its specific

Bergman called it olkali vegetabile vitriolotum, and Motveau vitriol gotarb.

gravity according to Wallerius " is 2'298; scome Hassenfratz, 2 4073 †; according to Watson, re-At the temperature of 60° it dissolves in 16 in weight of water; in a boiling heat it is soluble times Its weight s. When it is boiled in water liquid dissolves 0.242 parts, or nearly 4th of its of it II.

It suffers no alteration when exposed to the zir. placed upon burning coals, it decreptates, but side other alteration. In a red heat it melts, and imper cent. of its weight. It is sometimes luminous dark, as Mr Giobert has observed T.

Its component parts, according to the analyses to published, are as follows:

		++	- 55	İt	Į III	1
Acid	36-4	38 5	40	42.2	45'2	ė,
Base	63*6	61.5	52	58.1	54.8	11
Water			g	7.7		
Total	100.0	100	100	100	100	1

Variety 2. Supersulphate. This salt may be preby heating together a mixture of three parss of sul

[&]amp; Cormistry, p. 265.

Ann. de Chim. 23viii.

I Chemical Erroys, V 67.

[§] Bergman, i. 135

Wenze 's Vermandticheft, p. 320.

⁹ Ano. de Chien, 2, 40.

[&]quot; Thenard and Roard, Ann. de Côin. lit. 69.

⁺⁺ Rick

⁶ Bergman, i. 233.

It This is the result of my analysis. 200 Grains of fused miles porash gave 128 of sulphate of barytes. If this contains 35 per al áud, it will indicate 42'3.

B Kirwan, Nicholson's Jour. iii. 215.

potash and one part of sulphuric acid. It was first distinguished from the first variety by Rouelle senior.

Chap. III.

It usually crystallizes in long slender needles, but it Proporties may be obtained in six-sided prisms. Its taste is acrid, and it reddens vegetable blues. It is soluble in five parts of water at the temperature of 6000. It is scarcey altered by exposure to the air. When heated it melts readily and assumes the appearance of oil, but it becomes as white as ever when allowed to cool. A very violent and long continued heat dissipates the exmess of acid, and converts it into sulphate. Dr Wollaston has shown, that this salt contains just twice as much acid as the sulphate +.

But besides these two varieties, there are several others not yet particularly described. At least I have sound the proportion of acid to vary considerably in difberent states of the salt.

Sp. 2. Sulphate of Soda.

This salt was first discovered by Glauber, a German chemist, and for that reason was long known by the name of Glauber's salt. He himself called it sal mira-Mile. It may be prepared by saturating soda with sulphuric acid; but it is more usually obtained by dea composing common salt in order to procure muriatic acid. Like the sulphate of potash, it is capable of existing in two states. In the one it is neutral, in the other at has an excess of acid.

Variety 1. Sulphate. This is the state in which the

Link, Crell's Annals, 1796, i. 26.

⁴⁻On experacid and subscid sales. Phil. Trans. 1808.

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Properties.

great quanties by the manufacturers of sal amne a list crystals are transparent, and when formed by an evaporation, are six sided prisms, terminated by dilectal summits. The sides of the prisms are usually annuelled, and the crystals are almost always exceeding irregular. Its specific gravity, according to Walliam is 2.246°; Dr Watson, by a method not suscepted much precision (on account of the ready solubility this salt), found it 1.350 †.

It is soluble in 2.86 times its weight of water at temperature of 60°, and in 0.8 of boiling water 1. Whe heated to redness, so as to drive off its water of tristalization, and then pounded in a mortar, it dissolves it 3.3 its weight of water at the temperature of 144° When exposed to the air, it loses great part of its mater, and falls into a white powder [], but it is not oder wise altered. It loses about 56 per cent. of its weight

When exposed to heat, it first undergoes the water fusion **, then its water is evaporated, it is reduced a white powder, and in a red heat it melts. It kirwan has observed, that part of the acid, as well the water, is driven off by the application of a strotheat ††.

Wallerius' Chemia Physica, 266.

^{*} Ghamical Essays, v 66.

[|] Bergman, k 134.

Wenzel, p. 109

A Substances which fail to powder in this manner are said to garage

Wenzel, p. 312.

water they contain, they are said to undergo the watery fusion.

[†] Irisb Transactions. V.

Its constituents, according to the most accurate ann.

Chap. III.

	* _	+	‡	ğ
Acid	23.52	27	55*7	56
Base	18:48	15	44.3	44
Water	58.00	58		
Total	100	100	100	100

These analyses differ but little from each other; that of Kirwan is probably the nearest precision.

Variety 2. Supersulphate. When the sulphate of soda is dissolved in sulphuric acid, if the solution be set aside, it deposites spontaneously large rhomboidal crystals which contain an excess of acid. They effloresce in the air, and lose their excess of acid when moderately heated. At the temperature of 66° they dissolve in twice their weight of water ||.

Sp. 3. Sulphate of Ammonia.

This salt was discovered by Glauber, and called by him secret sal ammoniac. It was also called vitriolated ammoniac. It may be prepared by saturating ammonia with sulphuric acid, or by decomposing sal ammoniac by means of sulphuric acid.

^{*} Kirwan, Nicholson's Quarte Jour. 16. 215.

⁺ Bergman, 1. 133. | Wenzel's Verwondtschaft, p. 56.

Kirwan, Ibid. The two first analyses were made upon the crystals of the sait, the two last upon the sait supposed free from water | Link, Crell's Annals, 1796. i. 27.

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Propertues.

Its crystals are generally small six-sided prisal planes are unequal, terminated by a x-sided political later at the temperature of 60 its own weight of water at the temperature of 60 its own weight of boiling water. Water of the rature of 1440 dissolves 0.78 of its weight of the When exposed to the air, it slowly attracts made

When heated, it first decrepitates, then meliclose vessels sublimes, but with some loss of lit. In that state it might be called superman ammonia, as it contains an excess of acid, has taste, and reddens vegetable blues. When heat ly to redness, the greatest part of it is decresome sulphurous acid comes over and some a great quantity of azotic gas is disengaged, a probability water is formed 1.

Its constituents have been determined as fol

Composi-

			1
Acid	54160	58:75	59.8
Base	14.24	41.25	40.2
Water	31.10		
Total	100	100	100

Wennel, p. 309. † Kirwan's Min. is. 22

[§] Hatchett, Phd. Trong. lanavi. 314. § Kirwan, Nicholson's Suzzto Jose. iii. 215.

Wentel . Ferwandtrchaft, p. 58.

A Richter, Stategne Chemique, 1. 236.

These analyses differ very much from each other in the proportion of ammonia. The statement of Kirwan appears to be the most correct.

Sp. 4. Sulphate of Magnesia.

This salt is held in solution in the springs at Epsom in England, and was procured from them by evaporation more than a century ago. Hence the term Epsom salt, by which it was long distinguished. Some account of it was given by Grew in 1675; and in 1723, Mr Brown published a description of the process employed in extracting it from the springs, and in purifying it . In Italy it is manufactured from shistose minerals, containing sulphur and magnesia. By roasting these minerals, and then moistening them and exposing them to the air, the salt effloresces on their surface. By solution in water, with the addition of a little lime to precipitate any metallic substance that may be in solution, and repeated crystallizations, the salt is obtained in a state of purity +.

It exists in considerable quantity in sea-water; and the uncrystallized residuum in the salt pans, after all the common salt has crystallized, consists partly of this galt dissolved in water. This residuum is usually called bittern, and sometimes in Scotland spirit of salt.

Sulphate of magnesia crystallizes in quadrangular Properties. prisms, whose planes are equal, surmounted by quadrangular pyramids, or by dihedral summits. primitive form of its crystals is, according to Hauy, a four-sided prism, whose bases are squares. The form of

[•] Phil. Trans. 22211. 348.

Ann. de Chim. alviit. 80. Sec also Gehlen's Jour. ili. 549.

Total K

the integrant particles is a three-sided prior, where he see the toring interest interest in a standard prior of the cycle and desired prior interest. The cycle are desired prior interest.

It is an intervely bitter water. Its specific grain is 1714. At the temperature of 1819 it is sainble a in own weight of water, and in loss than two-thirds of its weight of boiling water? The volume of wat i increased faths by assing the sait I. When experit to the sirk effective, and is mourant to powder. The expected to hear a unsergoes the watery fusion; and y increasing the temperature its water to exaporated, by it cannot be decomposed by means of hear. Before it blow-pipe it meles with difficulty into an against vitatus globale 11.

The following Table exhibits the sessit of the experiments of different chemists to determine the properties of the constituents of this said.

Compad-

	1 5	1	4	60 :	**	; 5
Acid .	29-35	30-64	32	33	61-9	63-32
Base	17	16.86	19	19	35-1	36.68
Water	53.05	52.20	49	48		
Total	100	100	100	100	100	100

⁴⁴ Traité de Mineralogie, ii. 322.

^{*} l'amenfratz, Ann. de Chim. 22 viii. 12. † Bergman, Opac. i. 135.

Kirwan, Nicholenn's Querte Jour. iii. 215.] Wenzel, p. 62

⁹ M. J. Ann. de Chim. xlviii. 81. 4# Bergman, i. 135.

⁴⁴ Michter, Statlyne Chemique, 1. 136.

These analyses do not differ much from each other; probably a mean of the whole would come pretty near the accurate result.

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Sp. 5. Sulphate of Potash-and-Ammonia.

This salt, which was first described by Link, may be formed by saturating the supersulphate of potash with ammonia. The crystals are brilliant plates, having a bitter taste, and not altered by exposure to the air. They are composed of

60 sulphate of potash 40 sulphate of ammonia

100 *

Sp. 6. Sulphate of Potarb-and-Magnesia.

Link. He formed it by saturating supersulphate of potash with magnesia. Berthollet formed the same salt by mixing together equal solutions of sulphate of potash and muriate of magnesia, and evaporating the mixture. Crystals of sulphate of potash, with a little muriate of magnesia, are first obtained, and then the triple salt in rhomboidal crystals, which are not altered by exposure to the air. Their solubility is nearly the same as that of sulphate of potash †.

The taste of this salt is bitter. It is composed of

- 3 parts sulphate of potash
- 4 parts sulphate of magnesia

71

† Mem. de l' Instit. iii. 218.

^{*} Link, Crell's Annals, 1796. i. 19. Link, Crell's Annals, 1796, i. 30.

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Sp. 7. Sulphate of Soda-and-Ammonia.

This salt was first described by Link, who formed a by saturating supersulphate of soda with ammonia. See guin formed it by mixing together the solutions of inhalter of soda and sulphate of ammonia, and evaporating the mixture. The triple salt is obtained in crystals. These crystals are regular, and are not altered by especture to the air. Their taste is pungent and button. When heated they decrepitate and swell, ammonia and solutions discongaged, and supersulphate of ammonia and solutions of soda remain. Soda decomposes it by driving off the ammonia.

According to Link it is composed of

5 parts sulphate of sods

o parts sulphate of ammonia

14+

Sp. 8. Sulphate of Soda-and-Magnesia.

LINK obtained this salt by saturating supersulphate of soda with magnesia, and evaporating the solution. The crystals are presentic, large, have a bitter taste, and effloresce in the air. It is composed of

5 parts sulphate of soda

6 parts sulphate of magnesia

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♥ Jur. de Min. An. 20. p. 80.

† Crell's Annals, 1796, i. 34.

\$ Link, Ibid.

86. 9. Sulphate of Magnesia-and-Ammonia.

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This salt was pointed out by Bergman; but Fourcroy was the first chemist who examined its properties §. It may be prepared by mixing together saturated solutions of sulphate of ammonia and of magnesia. Crystals almost immediately precipitate, which consist of the two salts combined together. Or it may be prepared by pouring ammonia into a solution of sulphate of magnesia. Part only of the magnesia is precipitated. This is to be removed by filtration; and on evaporating the solution, the compound salt is obtained in crystals.

This salt crystallizes in octahedrons. Its taste is acrid and bitter. Its specific gravity is 1.690°. It is less soluble in water than either of its component parts. When heated it undergoes the watery fusion; and if the heat be increased, it is decomposed. It is not altered by exposure to the air.

It is composed, according to the analysis of Four-

Compad-

68 sulphate of magnesia 32 sulphate of ammonia

100

Sp. 10. Sulphate of Alumina.

This salt may be formed by dissolving alumina in sulphuric acid, evaporating the solution to dryness, dissolving it again in water, and evaporating it till it crystallizes. Little attention has hitherto been paid to this salt, which was never properly distinguished from alum

[·] Hassenfrags, Ibid. zaviji. 231.

Book II. Divinon III. till two memoirs, one by Vanquelin and more a Chaptal, on the nature of alum, made their appeared in the 22d Volume of the Annales de Chinic.

Properties.

It crystallizes in thin plates, soft and plant, and at pearly lustre. Its taste is astringent. It is very some in water, and does not crystallize without different When heated it loses its water of crystallization, and falls to powder. A strong heat decomposes it completely by volatilizing the acid. It is not altered a exposure to the air.

Sp. 11. Alum.

Varieties.

Or this salt there are no less than four varieties, if of which are triple salts; two neutral, and two in the state of supersalts. These varieties may be distinguished by the following names:

- 1. Sulphate of alumina-and-potash.
- 2. Sulphate of alumina-and-ammonia.
- 3. Supersulphate of alumina-and-potash.
- 4. Supersulphate of alumina-and-ammonia.

The two last of these varieties are usually confound ed together under the name of alum: the two first have been called alum saturated with its earths, and some times aluminated alum. It will be proper to consider the two last varieties first, because they have been longest known, and are the most important.

Variety 1. The supersulphates. The everyon of the Greeks, and the alumen of the Romans, was a native substance, which appears to have been nearly related to green vitriol or sulphate of iron; and which consequently was very different from what we at present denominate alum. From the researches of Protessor Beckman, it appears that we owe the discovery of alum

the discovery was made, is altogether unknown. It continued to be imported from the east till the 15th century, when a number of alum works were established in Italy. In the 16th century it was manufactured in Germany and Spain; and during Queen Elizabeth's reign an alum work was established in England by Thomas Chalomer. The alum of commerce is usually obtained from native mixtures of pyrites and clay, or sulphuric acid and clay.

Bergman has published a very complete dissertation on the process usually followed. The earth from which it is procured is usually called aluminous schistus, because it is slaty. Its colour is blackish, because it contains some bitumen. In most cases it is necessary to burn it before it can be employed; this is done by means of a slow smothered fire. Sometimes long exposure to the weather is sufficient to produce an efflorescence of alum on the surface. It is then lixiviated, and the water concentrated by evaporation, and mixed with putrid urine, or muriate of potash; crystals of alum and of sulphate of iron usually form together.

The composition of alum has been but lately understood with accuracy. It has been long known, indeed, that one of its ingredients is sulphuric acid; and the experiments of Pott and Margraf proved incontestibly that alumina is another ingredient. But sulphuric acid and alumina are incapable of forming alum. Manufac-

[·] Opusc. i. 279.

[†] Some chemises have thought proper to call the sulphuric acid, ubteined by distrilling alum, spirit of alum.

Book II. DivisionIII. or of ammonia, or of some substance containing the alkalies, is almost always necessary; and it was properthat in every case in which such additions are unnecessary, the earth from which the alum is obtained contained already a quantity of potash. Various conjectures a made about the part which potash acts in this case, Vauquelin and Chaptal † appear to have been the chemists that ascertained by decisive experiments to alum is a triple salt, composed of sulphuric acid, aluma, and potash or ammonia, united together.

Properties.

Alum crystallizes in regular octahedrous, consistent of two four-sided pyramids applied base to base. To sides are equilateral triangles. The form of its in grant particles, according to Hany, is the regular telehedron. Its taste is sweensh and very astringent. always reddens vegetable blues. Its specific gravity 1.7109 1. At the temperature of 00° it is soluble from 15 to 20 parts of water, and in 1ths of its weight of boiling water. When exposed to the air it die resces slightly. When exposed to a gentle heat it me dergoes the watery fusion. A strong heat causes it swell and foam, and to lose about 44 per cent. of in weight, consisting chiefly of water of crystallizations What remains is called calcined or burnt alum, and sometimes used as a corrosive. By a violent heat, the greater part of the soid may be driven off. In that can as was first observed by Gay-Lussac, a portion of

[.] Ann. de Chim. xxii. 258. ∮ 1bid. 285.

[†] Hamenfratz, Ann. de Chim. xxviii 12. Wallorsus Sound it 17 (Chemistry, p. 266); and De Watson, 1-757 Lecays, v. 67); Fahrent 2782. (Phil. Trens. liii. 114) § Bergman, i. 287.

ough the properties of alum are in all cases pretty the same, it has been demonstrated by Vauquenet three varieties of it occur in commerce. The
is supersulphate of alumina-and-potash; the sesupersulphate of alumina-and-ammonia; the third
mixture or combination of these two, and contains
potash and ammonia. It is the most common of
doubtless, because the alum makers use both urine
muriate of potash to crystallize their alum. Vauin, Thenard, and Roard have lately analysed a numof specimens of alum manufactured in different
tries. The result was that they all contain very
ty the same proportion of ingredients. The mean
I their trials was as follows:

ч	Dap.	ш.
1		_
v.		
4	1.44-23	

	+	
Acid	30-52	25.04
Alumina	10.20	12.53
Potash	10-40	10.05
Water	48-58	51'41
Total	100-00	100.00

com the late analyses and experiments of Thenard

Small quantities of ammenia also fed in several

menard and Roard, Ann. & Chm. lix. 72. The seid was estimated insign sulphate of harytes, which they considered as containing only more of sulphuric send; whereas it contains 33 per cert; of source portion of that seed in the table is too small.

74 SALTS.

Book U. Division II. and Roard, it appears that alum usually contains a sulphate of iron. Its good qualities as a mordal dyeing, depend upon the proportion of that sate sent. The freer it is from it the better. The palum examined contained about when the sulphate of iron, the impurest about weight of sulphate of iron, the impurest about a weight of reed from sulphate of iron every specialism tried acts exactly in the same manner as a dant.

Subte alum.

When an unusual quantity of potash is added to liquor, the salt loses its usual form and crystaling cubes. This constitutes a fourth variety of alumnally distinguished by the name of cubic olumnations an excess of alkali.

Fifth va-

When the potash is still further increased, Chas observed that the salt loses the property of callizing altogether, and falls down in flakes. This stitutes a fifth variety of alum, consisting of sulph potash combined with a small proportion of alumin

Neutral compounds Variety 2. The sulphates. All the varieties of are capable of combining with an additional dose of mina, and forming pertectly neutral compounds possessed of nearly the same properties, and then confounded together as one salt.

Properties.

It may be prepared by boiling a solution of with pure alumns; the compound, as it forms, go ally precipitates in the form of a white powder. It formerly denominated alum saturated with its configuration of a tasteless powder. It is insoluble in water, and



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Itered by exposure to the air. Heat has no effect upon * t unless it be very violent; and in that case part of the ecid is disengaged. Sulphuric acid converts it into lum. It is less easily decomposed by other bodies than lum. Several of the acids dissolve slowly its last dose of alumina, and convert it into alum. This salt has not hitherto been applied to any use. Alum is capable also, as Chaptal informs us, of combining with several nother bases, and of forming many triple salts, which have never yet been examined with attention .

If three parts of alum and one of flour or sugar be Homberg's melted together in an iron ladle, and the mixture dried till it becomes blackish and ceases to swell; if it be then pounded small, put into a glass phial, and placed in a sand-bath till a blue flame issues from the mouth of the phial, and after burning for a minute or two be allowed to cool +, a substance is obtained known by the name of Homberg's pyrophorus, which has the property of catching fire whenever it is exposed to the open air, especially if the air be moist.

This substance was accidentally discovered by Homberg about the beginning of the 18th century, while he was engaged in his experiments on the human fæces. He had distilled a mixture of human fæces and alum till he could obtain nothing more from it by means of heat; and four or five days after, while he was taking the residuum out of the retort, he was surprised to see it take fire spontaneously. Soon after, Lemery the Younger discovered that honey, sugar, flour, or almost any

Paropherus.

Ann. de Chim. xxii. 293.

Care-must be taken not to keep it too long exposed to the heat.

Book II. Disseion III. animal or vegetable matter, could be substituted human faces; and afterwards Mr Lejoy de Savishowed that several other salts containing sulphacid may be substituted for alum. Scheele pathat alum deprived of potash is incapable of for pyrophorus, and that sulphate of potash may be a tuted for alum. And Mr Proust has shown another of neutral salts, composed of vegetable and earths, when distilled by a strong fire in a leaves a residuum which takes fire spontaneously posure to the air.

These facts have thrown a great deal of light anature of Homberg's pyrophorus, and enabled some measure to account for its spontaneous infation. It has been ascertained, that part of the aric acid is decomposed during the formation of the rophorus, and of course a part of the alkaline becomes uncombined with acid; and the charcoal, gives it its black colour, is evidently divided into minute particles. It has been ascertained, that the combustion of the pyrophorus a quantity of a is absorbed. The inflammation is probably occar by the charcoal; the sulphuret of potash also a essential part. Perhaps it produces a sudden in of temperature by the absorption and solidifications water from the atmosphere.

[·] See Macquer's Dietlenory.

[†] Scheele on Fire and on Pyresteria.

Sp. 12. Sulphate of Titria.

Chap. III.

THIS salt was first formed by Gadolin, and afterwards it was examined with more precision by Eckeberg , Vauquelin t, and Klaproth 1.

Sulphuric acid dissolves yttria readily, and some caloric is evolved during the process. As the solution goes on, the sulphate crystallizes in small brilliant grains.

These crystals are irregular, but most frequently, Properties. according to Eckeberg, they have the form of flat sixsided prisms terminated by four-sided summits. These crystals are not altered by exposure to the air. Their taste is astringent and sweetish, though not so much so as sulphate of glucina. Their colour is a light amethyst red; their specific gravity 2.791. They require at least 30 parts of water at the temperature of 60° to dissolve them. At a red heat they are partly decomposed.

Oxalic acid, prussiate of potash, and the infusion of outgalls, occasion a precipitate in the aqueous solution of this salt. It is decomposed by phosphate of soda.

Sulphate of glucina is readily distinguished from this salt by its being colourless, lighter, and more soluble in water.

Sp. 13. Sulphate of Glucina.

This salt was first examined by Vanquelin, the discoverer of its base.

Beitrage, in. 67.

Creil's Annals, 1799, ii. 68.

[†] don de Chima unvi. 156

Book I'. Division III. It is prepared by saturating sulphuric acid with given a. The acid dissolves the earth readily, and the lution by evaporation yields small needle-form cryst the figure of which has not been ascertained.

Properties.

It is very soluble in water; the solution readily assumes the consistence of a syrup, but is brought and difficulty to crystallize. When heated it undergoest watery fusion, loses its water of crystallization, and to powder. At a red heat it is decomposed complete the acid is driven off, and the earth remains in a solution of this salt, occasions a yellowish who precipitate.

Sp. 14. Sulphate of Zirconia.

This sait was first formed by Klaproth, but Vauque lin has described it with more precision. It is preparably dissolving zirconia in sulphuric acid, and evaporating the solution to dryness.

Propert'es.

however, be obtained in small needle-form crystals. has no taste, and is insoluble in water. It is not altered by exposure to the air. It is very easily decomposition by heat: The acid readily shes off, and leaves the entire in a state of purity. This decomposition may be accomplished even by boiling in water; the earth precipitates, and the acid remains in solution. Klaproth is forms us, that with excess of acid sulphate of zirconforms transparent stelliform crystals, soluble in water and having an astringent taste.

[#] Joer. de Phys. 22241. 287.

Chap. III.

II. IN SOLUBLE SULPHATES.

Sp. 15. Sulphate of Lime.

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Of this salt there are two varieties. The first contains water, and is called common sulphate: The second, which is destitute of water, is called anhydrous sulphate.

Variety 1. Common sulphate. This salt was well known to the ancients under the name of gypsum; but the composition of gypsum was not known till Margraf and Macquer analysed it, and proved it to be a compound of sulphuric acid and lime. The salt formed by the artificial union of these two bodies was formerly called selenite, probably from its whiteness.

The properties of this salt were first examined with Properties. precision by Bergman. It is found abundantly in different parts of the world, so that it is seldom formed artificially. When pure, it is frequently crystallized. The primitive form of its crystals is, according to Hauy, a right angular prism with rhomboidal bases, whose angles are 113° and 67°. Its integrant particles have the same form; but it is more usually found crystallized in octahedrons, in six-sided prisms, with four-sided summits, or lenticular. These crystals are often exceedingly transparent.

It has a slightly nauseous taste, scarcely perceptible, except by drinking a glass of water impregnated with it *. It is soluble in 400 parts of water at the temperature of 60° †, and in 450 of boiling water. Mr Paul has observed, that if this solution be saturated by pres-

+ Bucholz, Gehlen's Just. v. 165.

Book II. Division III. sure with hydrogen gas, the sulphate in about sire is converted into a sulphuret ‡‡. The salt is not by exposure to the air. It is soluble in sulphate

When heated, it loses its water of crystallizate crepitates, and falls into a soft white powder powder, when its water has been driven off by plication of a red heat, absorbs water very rapit solidifies it; at the same time a slight increase perature takes place: so that if it be formed into with water, it dries in a few minutes. In this is called plaster of Paris, and is much empired forming casts, &cc. in consequence of this proper

Sulphate of lime, when exposed to a viole melts; before the blow-pipe it gives an opaquous globule. The température necessary to this effect is, according to Saussure, 51° of wood §§.

The following Table exhibits the result of the ent experiments hitherto made to aspertain the tion of the constituents of this salt.

Composi-

Î		H	H	‡	9			**	1
ı	Acid	43	43	4 b	55.75	56:58	57	57.57	Ľ.
ı	Base	33	57	32	44.25	43-42	43	42.49	1
į	Water	24	À	22					
i	Total	100	100	100	100	100	100	100	100

^{\$1} Phel. Mag. zv 63.

ji Jaw. de Phys. 210 Ohenevix, Phat Ma

The third is

Bucholz, Gehlen's Jur. v. 162.

[#] Hergman, i. 135. By my own analysis.

Kirwan, Nicholson's Querte Jour

These numbers, except the analysis of Chenevix, do not differ much from each other. The analysis of Bucholz seems most correct. It scarcely differs from the result of my experiments.

Chap. III.

Anhydrous Sulphate.

Variety 2. Anhydrous Sulphate. This variety is found native in different parts of the earth; Sweden, Tyrol, Berne, &c. It seems to have been first pointed out by Hauy. A specimen of it was described with much precision by Fleuriau in the Journal de Phisique for 1798; and analysed by Vauquelin, who first ascertained its composition. It has since been described with more precision by Bournon; and a purer specimen of it analysed by Chenevix; and more lately by Klaproth §.

It is usually crystallized. The primitive form of its crystals is a rectangular prism, having two of its faces broader than the other two. It has considerable lustre, and the broad faces have the appearance of pearl. Its specific gravity is about 2.950. Its hardness is considerable, being not inferior to that of calcareous spar. It usually phosphoresces when heated. It is transparent, insoluble in water, and in its other properties agrees with common sulphate.

In the proportions of its constituent parts it coincides exactly with common sulphate, excepting in the absence of water.

Sp. 16. Sulphate of Barytes.

This salt is found abundantly in different parts of the earth, and was formerly denominated ponderous spar,

Jour, de Min. An. z. ii. 345.

^{\$} Ibid. p. 418,

Gehlen's Jour. ii, 355.

Book H. Division H.L. from its great weight. Its composition was fire to tamed by Gabo.

I'd parties.

Hany has shown that the form of its integrant ticles is a right-angled prism, whose bases are remained with angles of 10110 and 7610. The primitive to of its crystals is the same, but it occurs native in an riety of other forms. It cannot be crystallized by This salt is insoluble in water, or at least only sold in 43000 times its weight of water at the temperal of the atmosphere. Sulphuric acid dissolves it is concentrated and boiling, but it is precipitated by addition of water 1.

When suddenly heated, it breaks in pieces and about with a crackling noise. This phenomenon, whi is called decrepitation, is occasioned by the sudden enversion of the water which it contains into stee When heated very violently it melts, and before to blow-pipe is converted into a white opaque global According to Saussure, a heat equal to 35" Wedgene is required to fuse it 1. When formed into a thin call with flour and water, and heated to redness, it phospheresces in the dark. This was first observed in a variety of this substance known by the name of Bologna stoe Lemery informs us, that the property was discoverably an Italian shoemaker named Vincenzo Casciard

Kirwai, & Mor. i 136.

With ring first observed. Mr Hume has ano mentioned it in Phil Many and the face was well known to chemists. This solution, he ever, is but very imperfectly critical to the name of supersulphate baryees; is it parts with its exceedul acid at a moderate heat, which has no and is the case with a perfect of potash.

¹ Jear, de Phys. siv. 15.

This man found a Bologna stone at the foot of Mount Paterno, and its brightness and gravity made him suppose that it contained silver. Having exposed it to the fire, doubtless in order to extract from it the precious metal, he observed that it was luminous in the dark. Struck with the discovery, he repeated the experiment, and it constantly succeeded with him. It is evident that by the calcination it must be converted, at least partly, into sulphuret.

The following Table exhibits the result of the different experiments hitherto made to ascertain the composition of this salt.

	†	‡	}	11	9	**	++
Acid	24	25.18	31	32	33	33.96	34
Base	76	74.82	69	68	67	66.04	бб
Total	100	100	100	100	100	100	100

Composi- . -

These results differ very considerably from each other. The analysis of Kirwan seems most correct. It is of great importance to have an exact analysis of this salt, as it is by means of it that the proportion of sulphuric acid in different compounds is determined.

[†] Chenevix, Nicholsou's Jour. ii. 196.

[†] Thenard, Ann. de Chim. xxxii. 266.

[§] Bucholz, Beitrage, iii. 31.

[|] Vauquelin, Ann. de Chim. 1, 168.

[¶] Kirwan and Kiaproth, Gehlen, v. 515.

^{**} Aiken, Nicholson's Jour. xxii. 304.

[#] Fourcroy, iii. 25.

Book II. Division III.

Sp. 17. Sulphate of Strontian.

This salt, for the discovery of which we are indeed to Dr Hope and Mr Klaproth, may be composed a tificially by dropping sulphuric acid into strontian a ter; in which case it has the form of a white powd-But it exists abundantly in different parts of the worusually crystallized in rhomboidal prisms.

Properties.

This salt is tasteless. It is soluble in 3640 parts boiling water. Sulphuric acid dissolves it readily we assisted by heat, but it is precipitated by the addition water to the solution. In the greater number its properties it has a considerable resemblance to sphate of barytes.

Contpool-

It is composed, according to the analysis of Vaug

54 strontian

100

58 strontian

100

This salt is soluble in sulphuric acid, and thereforens an imperfect supersulphate.

[·] Hope, Teans. Edin. iv. 10.

^{\$} Bestrage, 11, 97.

f lbid p. #37.

^{**} Hume, Phil. Mag. 214. 358.

⁺ Joor. de Min. Att. vi. 6.

Nicholaun's Journal, III. 1

^{†!} Ibid. p. 215.

Such are the properties of the sulphates. The following Table exhibits the solubility of each salt in water, and the proportion of the constituents, as far as ascertained. They are all insoluble in alcohol:

Sulphates of	Solubility in 100	Constituents.				
Julipunca St	Water.	Acid.	Bate.	Water.		
Ammonia	50	100	26.05	57		
Magnesia	100	100	57.92	182.8		
Lime	0.55	100	76.40	55.8		
Soda	35	100	78-32	246.6		
Potash	6.5	100	110	20		
Strontian	0.0	100	138			
Barytes	0.0	100	203			

THE SUL-

THE genus sulphates contains several salts of considerable importance.

1. Salphate of potash was formerly used in medicine; but on account of its disagreeable taste and little solubility it is now laid aside. Apothecaries, indeed, sometimes use it to facilitate the powdering of vegetable catharties. To the alum-makers it is of considerable importance, as it enters into the composition of alum, which cannot be obtained crystallized without potash or ammonia.

Book II. Division III.

- 2. Sulphate of soda is one of the most common of all the cooling purgatives, as they have been termed; immense quantities being annually manufactured at consumed.
- 3. Sulphate of ammonia is sometimes employed in chemical analyses. Thus it was used with advantage by Hatchett in examining the molybdate of lead.
- 4. As great quantities of the sulphate of magnet are prepared for commercial purposes, the salt is probably in common use on the continent as a purgative. Its intensely bitter taste has brought it into disuse this country. What is sold by a pothecaries under the name of Epsom salt is often nothing else than sulphate of soda in small irregular crystals.
- 5. Sulphate of lime is much used as a cement, under the name of plaster of Paris. It is employed also for moulds, and for casting into statues, &c. It is fine heated, in order to deprive it of its water. In this state it forms with water a liquid paste, which in a few minutes becomes solid in consequence of the combination of the water with the sulphate. It constitutes the charingredient of stucco; and is in many places used as a manure.
- 6. The sulphates of barytes and strontian are of applied to any useful purpose directly; but they furnish almost all the barytes and strontian which the chemists employ.
- 7. But the most important of all the sulphates is undoubtedly alum, which is applied by manufacturent to too many uses for a complete enumeration. To the dyer and calico printer this salt is indispensible, a alumina, its earthy base, forms one of the best of all the mordants, or substances which have the property of an

ing colouring matters in cloth. It is used by the tan- Chap. III. ners in the preparation of leather. Indeed, some kinds of leather may be considered as nothing else than skins altered by being steeped in a solution of alum in water. It is often very useful in clarifying liquors, from the property which alumina has of forming an insoluble precipitate with many vegetable substances. Surgeons sometimes take advantage of the properties of this salt, and use it as an astringent or styptic; but it would be too tedious to enumerate the purposes to which this salt is applied by paper-makers, goldsmiths, candle-makers, bookbinders, &c.

END OF THE SECOND VOLUME.

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